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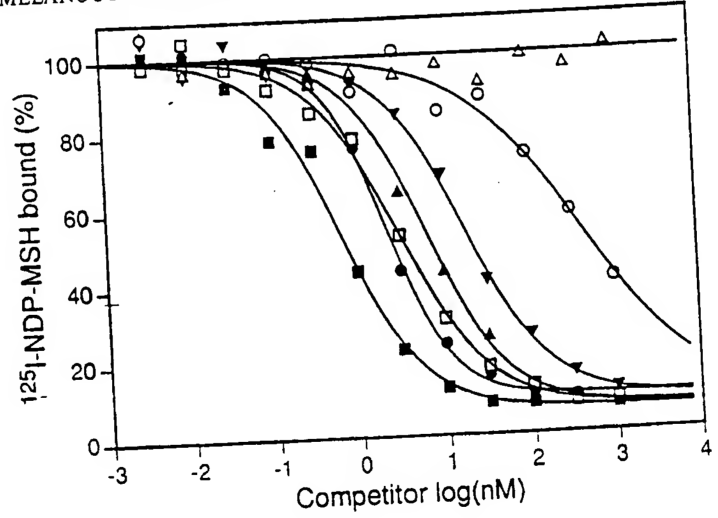
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(54) Title: HUMAN MELANOCYTE STIMULATING HORMONE RECEPTOR



Novel DNA fragments encoding novel polypeptides having properties of melanotropic hormone receptors, especially DNA fragments encoding melanocyte stimulating hormone receptors (MSH receptors), as well as polypeptides which are MSH DNA fragments encoding melanocyte stimulating hormone receptors (MSH receptors), as well as polypeptides for production of receptors, are disclosed. The use and engineering of melanotropic hormone receptor DNA and polypeptides for production of monoclonal antibodies for diagnostic and therapeutic purposes, as well as the engineering of drugs, cell lines, vectors and DNA monoclonal antibodies for diagnostic and therapeutic purposes, as well as the engineering of drugs, cell lines, vectors and DNA for therapeutic and diagnostic purposes are also disclosed. Also disclosed are methods for therapy and diagnosis of malignant for therapeutic and diagnostic purposes are also disclosed. Also disclosed are methods for therapy and diagnosis condition, impaired melanoma, skin cancer, vitiligo, pyretic condition, inflammatory condition, nociceptive condition, catatonic condition, impaired melanoma, skin cancer, vitiligo, pyretic condition, inflammatory condition, epilepsy and nerve damage, using the DNA fragmemory condition, reduced or increased skin tanning, pigmentation condition, epilepsy and nerve damage, using the DNA fragmemory condition, reduced or increased skin tanning, pigmentation condition, epilepsy and nerve damage, using the DNA fragmemory condition, polypeptides and antibodies. Methods for selecting substances which interact with the receptors are also disclosed.

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HUMAN MELANOCYTE STIMULATING HORMONE RECEPTOR.

#### FIELD OF THE INVENTION

The present invention relates to a DNA fragment encoding the human melanocyte stimulating hormone receptor (MSH receptor) 5 or an analogue or subsequence thereof. The DNA fragment contains an open reading frame of 951 bp which codes for a polypeptide of 317 amino acids, said DNA fragment as well as its analogues, subsequences or modifications constitute an important aspect of the invention. The DNA fragment has been 10 expressed in an eukaryotic cell line and the expressed protein has been found to have properties identical to that of a native MSH receptor. The invention also relates to a DNA fragment encoding a subtype of the human MSH receptor (in this application designated MC-2) which contains an open reading frame of 975 bp which codes for a polypeptide of 325 15 amino acids, said DNA fragment as well as its analogues, subsequences or modifications also constitute an important aspect of the invention. This DNA fragment has also been expressed in an eukaryotic cell line and the expressed protein has been found to have properties to that of an MSH receptor subtype.

The invention also relates to a polypeptide encoded by a DNA fragment of the invention and to analogues and subsequences of said polypeptide. Furthermore, the invention relates to the use of the DNA fragments or analogues or subsequences thereof, and to the use of polypeptides of the invention encoded by the DNA fragments of the invention. Especially interesting is the use of the polypeptides of the invention which have MSH receptor activity. The use of the polypeptides of the invention or analogues or subsequences thereof for generation of antibodies constitutes yet another aspect of the invention. Also, the invention relates to diagnostic and therapeutic methods and diagnostic and therapeutic agents for use in the diagnosis and treatment of MSH receptor expressing disease conditions such as vitiligo, melanoma, skin cancer,

pyretic conditions, inflammatory conditions and nociceptive conditions, catatonic conditions and impaired memory conditions, and to methods for detecting and quantitating the MSH receptor. In addition, the invention provides methods for 5 testing substances capable of interfering with the activity of the MSH receptor and methods for treatment of MSH receptor expressing disease conditions. The patent application also relates to the use of the MSH receptor coding fragments or the MSH receptor during non-disease conditions for the control or diagnosis and/or determination and/or production 10 control of skin and/or hair and/or fur colour in man and/or animals. Moreover, the patent application relates to the elucidation of the structure of the MSH receptor in three dimensions by the utilization of computer modelling methods 15 and/or by application of structure analysis by crystallographic approaches and/or NMR (Nuclear Magnetic Resonance) and to the use of the knowledge of the receptor structure for the design of drugs with binding affinity for the MSH receptor and/or its subtype (MC-2).

The present invention which comprises a DNA fragment encoding the MSH receptor or analogues thereof and the application of these and in this connection methods for identifying products which pertains to the MSH receptor and/or its biological functions constitutes significant contributions which will become useful for biotechnological, pharmaceutical, medical and veterinary practices. As a background to the uses of a DNA fragment and analogues and subsequences thereof and the application of these, some of the most important facts regarding the MSH receptor and its biological functions in man and animals are summarized below.

#### GENERAL BACKGROUND

Although information existed regarding the MSH receptor (reviewed below), the structure of the MSH receptor gene as well as the primary amino acid sequence of the MSH receptor has not been known before the priority date of the present

patent application. As appears from the following, the MSH receptor is a very important receptor with a number of different functions such as anti-inflammatory and antipyretic function and involved in a number of diseases such as melanoma and skin cancer and moreover, it is having an important role in the control of skin, hair and fur colour in man and animals.

MSH receptor and its biological functions

The MSH receptor belongs to a large class of receptors showing functional and structural similarities. These receptors 10 mediate their cellular effects via coupling proteins termed guanine nucleotide regulatory proteins (G-proteins), of which several types are known (e.g.  $G_{\text{s}}$ ,  $G_{\text{i}}$ ,  $G_{\text{k}}$  and others). The MSH receptor is a cell membrane bound protein which serves as a 15 recognition site for  $\alpha$ -MSH (melanocyte stimulating hormone). The term MSH relates to several peptides among which  $\alpha$ -MSH,  $\beta$ -MSH and  $\gamma$ -MSH may be mentioned; the  $\alpha$ -MSH generally showing the largest activity. These hormones are generally referred to as melanotropic hormones to which also the ACTH (adrenocorticotropic hormone) belongs as well as a number of related peptides, being present in man and animals. In the present patent application, peptides which have binding affinity for MSH receptors will collectively be referred to as MSH peptides or MSH receptor ligands. Upon binding of MSH receptor ligands to the MSH receptor, an activation of the receptor ensues which leads to altered activity of the cell in which the receptor is located. MSH receptors are known to be present in melanocytes which are pigment cells and in humans give the skin a varying amount of dark pigmentation and which have a role in protecting the skin from UV-radiation. In animals, melanocytes also have a role in skin pigmentation. In both animals and man changes in skin colour are at least partly mediated by melanocytes and these changes are also partly regulated by the degree of activation of MSH receptors by the peptide hormones that bind to the MSH receptor (Nord-35 lund 1991; Levine 1991).

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MSH receptors may also be localized in cell types other than melanocytes (Tatro 1987) where they may have other types of important physiological roles.  $\alpha\textsc{-MSH}$  is known to be produced in certain areas of the brain, such as the hypothalamus, 5 corpus amygdaloideum and cerebral cortex. Moreover, proopiomelanocortin, which is the precursor molecule for  $\alpha\text{-MSH}$ , is found in lymphocytes of the thymus and spleen, neutrophils, placenta, ovary as well as in the epidermis (Nordlund 1991). There is evidence that by acting on MSH receptors,  $\alpha\text{-MSH}$  may 10 have roles in (i) mediating neurotransmitter effects in the CNS, (ii) participating in endocrine regulation, (iii) modulating immune-inflammatory responses, besides (iv) regulating the skin pigmentation, as mentioned above (Nordlund 1991, Levine 1991). MSH receptors perform various functions in 15 neurochemical processes, such as the induction of antinociceptive effects, the perturbation of grooming behaviour, the alteration of stretch and yawn reflexes and the potentiation of catatonic states (Hirsh and O'Donohue 1986). Moreover, MSH receptors are implicated to have a function in the enhance-20 ment of visual and verbal learning (Veith et al. 1978; Ward et al. 1979, Handelman et al. 1983). The role for MSH receptors in endocrine function is indicated, for example, by observations that  $\alpha\textsc{-MSH}$  may affect cortisol secretion from the adrenal gland, and increase plasma levels of growth hormone, luteinizing hormone and follicle-stimulating hormone

MSH receptors also seem to be mediating the powerful antipyretic effect caused by  $\alpha$ -MSH (Clark et al. 1985) as well as the anti-inflammatory actions induced by  $\alpha$ -MSH (Rheins et al. 1989). Central MSH receptors are also involved in the mediation of anti-convulsive effects since MSH peptides exert anti-epileptic effects (De Wied 1993). Moreover, MSH receptors seem to mediate the growth factor effect of MSH peptides which mediates accelerated and enhanced nerve generation and muscle reinnervation after peripheral nerve injury (Strand et al. 1993).

(Reid et al. 1984).

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MSH receptor on melanoma cells

Melanoma cell lines are derived from immortalized melanocytes. Melanocytes are clinically the starting point of malignant melanoma (reviewed below). MSH receptors are present on many such melanoma cell lines, the reported frequency in different cell lines being more than 70% (Tatro et al. 1990a). In experimental melanoma, differentiation, tumorigenicity and metastatic potential of the melanoma are influenced by MSH (Kameyama et al. 1990). Moreover, α-MSH immunoreactivity has been demonstrated to be present in human melanoma metastases (Ghanem et al. 1989) indicating the possibility that locally formed melanotropic activity has a role in the pathogenesis of melanoma.

The presence of MSH receptors on melanoma cell lines suggests that endogenous  $\alpha$ -MSH, the major known form of circulating melanotropin in mammals, may modulate melanoma cell activity in vivo. The demonstration of specific binding sites in melanoma tumours does not prove that these are linked to cellular response systems in vivo, but this seems highly likely in view of the close relationship between binding and biological responses in cultured melanoma cells (Tatro et al. 1990b). Evidence suggests that  $\alpha$ -MSH may modulate proliferation and ability of melanoma cells to establish metastatic colonies (Lerner et al. 1989; Abdel-Malek et al. 1986).

It is well recognized that in mammalian melanocytes and melanoma cells α-MSH acts through MSH receptor on an intracellular pathway that involves the activation of adenylate cyclase (Tatro et al. 1990b). This leads to an increase in the production of cyclic AMP which in turn induces tyrosinase, a key enzyme in the melanin biosynthesis. However, there is evidence that melanotropins after binding to the MSH receptor increase the intracellular calcium (Mac Neil et al. 1990). It is conceivable that this effect is due to the fact that MSH/MSH receptor complex activates phospholipase C (PLC), which then acts to produce inositol 1,4,5-

trisphosphate, which then in turn triggers mobilization of intracellular calcium. This proposition is due to the fact that receptor mediated activation of PLC is a G-protein linked event, and that it has been shown that receptors may simultaneously, e.g. in a promiscuous way, act via several of the known G-protein linked metabolic pathways (Traiffort et al. 1992 and Gudermann et al. 1992). Activation of phospholipase C also leads to the production of diacylglycerol, the activator of protein kinase C. Indeed, it has recently been shown that MSH can activate protein kinase C (Buffey et al. 1992). Two other G-protein coupled receptors, namely the  $\alpha_{\text{1b}}$ adrenergic receptor and serotonin receptor, which are also coupled through the above mentioned second messenger system, are shown to be protooncogenic (Allen et al. 1991; Julius et al. 1989), thus further indicating the possibility that MSH receptor may have a pathogenic role in melanoma. Moreover, melanotropins are shown to induce expression of the growth associated oncogene c-fos (Hart et al. 1989) further supporting this notion. Note also Sukhanov et al. (1991).

#### 20 Malignant melanoma

Malignant melanoma (melanocarcinoma) is a malignancy derived from melanocytes. About 1% of all malignant tumours are malignant melanomas. The incidence of malignant melanoma is increasing rapidly. During the last decades the incidence has approximately doubled every 10 years with both sexes being affected equally. Malignant melanoma can develop at every site of the skin. There are sites of predilection: feet followed by head and neck. Infrequent sites are the genital organs, perineum, perianal region and mucous membranes. The tumour has a high incidence of metastasis to adjacent skin and regional lymph nodes. Haematogenous metastasis may also occur.

The main factor for the development of malignant melanoma is exposure to sunlight. The people who are mainly affected are those who have fair skin that can be easily damaged by the

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sunlight. Despite various therapeutic regimes the 5 year survival in melanoma with distant metastasis is only 5% and with regional metastasis it is 43% (Roses et al. 1991). Existing clinically approved therapies, besides surgical 5 removal of lesions, are non-specific and include limb perfusion, chemotherapy, immunotherapy, radiotherapy and hormonal therapy (Ho et al. 1990). Radiopharmaceuticals such as iodoquinoline (Lambrecht et al. 1984), iodothiouracil (Coderre et al. 1986) and N-(2-diethylaminoethyl)-4-iodobenzamide (Michelot et al. 1991) have been used for the diagnosis and therapy of melanoma, albeit with very limited success. Another approach for the diagnosis and therapy of melanoma is to use radiolabelled monoclonal antibodies against melanoma associated antigens (Eary et al. 1989; 15 Larson 1991). This poses the problem of having a true melanoma associated antigen. Also, different antigens are expressed based on the developmental stage of the melanoma tumour, and different tumour sites in the body may be expressing different antigens. This would require the use of a 20 mixture of monoclonal antibodies, all of them with very high specificity. The composition of such a mixture will vary between patients and between different tumour stages of the same patient. All this would be very difficult to achieve. Previous work has shown that MSH receptors are detectable in melanoma metastases of about 80% of human patients (see Tatro 25 et al. 1992).

MSH receptors on the melanoma cells have been considered as potential targets for novel drugs useful for treatment of the disease. Diphtheria toxin and α-MSH fusion protein have been constructed and shown to be selectively toxic for MSH receptor bearing cells in vitro by a targeted delivery of the diphtheria toxin (Murphy et al. 1986; Wen et al. 1991; Tatro et al. 1992). In another approach MSH was coupled with an antibody directed towards the CD3 receptor of cytotoxic T cells. The complex was shown to mediate cell lysis of melanoma cells in vitro. The MSH moiety binds to the MSH receptor of the melanoma cells whereas the antibody tags CD3 bearing

cytotoxic T-cells which mediate lysis of the melanoma cell (Liu et al. 1988).

MSH receptor and skin tanning and control of hair and fur colour

Endogenous and exogenous melanotropins are suggested to enhance human cutaneous pigmentation in vivo (Levine 1991; Mulligan et al. 1982; Lerener et al. 1961). The mechanism of action by which MSH and other melanotropins stimulate melanogenesis is well studied. The melanotropins bind MSH receptors on melanocytes and result in the activation of adenylate cyclase. Increased cAMP activates tyrosinase enzyme which converts tyrosine to dopa and dopa to dopaquinone, resulting in melanin formation. The melanin thus formed is partly secreted from the melanocytes and taken up by keratinocytes of the skin thus making the skin colour become more dark. Moreover, the pigment thus formed will constitute the colour of hair and fur in man and animal. Various colours will be produced depending on the level of the presence of melanin pigment in the hair, fur and skin.

MSH receptor and anti-pyretic and anti-inflammatory actions

The  $\alpha$ -MSH is one of the most potent antipyretic agents identified (Clark et al. 1985). Moreover, both afferent and efferent inflammatory responses to chemicals and irritants, like phorbol esters or contact allergens, are blocked by the topical application of  $\alpha$ -MSH. These anti-pyretic and anti-inflammatory effects seem to reside in the carboxy terminal region of the hormone supporting the notion of their mediation via an action on MSH receptors.

### 30 MSH receptor and vitiligo

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In vitiligo areas of loss of skin pigmentation is a characteristic feature. Such loss of skin colour is due to loss

and/or malfunction of pigment cells. Due to localization of MSH receptor on skin pigment cells it is considered that MSH receptor has a role in vitiligo. One of the causes of the vitiligo can be autoimmune reactions of the host against the MSH receptor protein and/or polypeptides. Thus, the MSH receptor constitute an interesting target in the cure and/or amelioration of the vitiligo condition.

#### DISCLOSURE OF THE INVENTION

The above summarized activity and involvement of the MSH receptor in a number of biological functions of various cells clearly shows the importance of the present invention which relates to a DNA fragment encoding a polypeptide having MSH receptor activity. Despite considerable efforts to elucidate the sequence of such a DNA fragment, nobody had prior to the present invention succeeded in doing this.

Accordingly, the present invention relates to a novel DNA fragment having the nucleotide sequence shown in SEQ ID NO: 1 or an analogue or subsequence thereof which

- 1) has a homology with the DNA sequence shown in SEQ ID NO: 1 of at least 50%, and/or
  - 2) encodes a polypeptide, the amino acid sequence of which is at least 50% homologous with the amino acid sequence shown in SEQ ID NO: 2, and/or
- 3) encodes a polypeptide which binds an antibody which is also bound by an MSH receptor, and/or
  - 4) encodes a polypeptide which is an MSH receptor or which has the same binding capacity as an MSH receptor.

The DNA fragment with the nucleotide sequence shown in SEQ ID NO: 1 is derived from a human cDNA library and has been found

to contain an open reading frame of 951 bp which codes for a previously unknown polypeptide of 317 amino acids which is shown in SEQ ID NO: 2. This polypeptide constitutes the entire polypeptide of an MSH receptor.

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A detailed description of the molecular cloning and nucleotide sequencing of the cDNA on the basis of the carefully constructed primers is given in Example 1. The cDNA of the MSH receptor represents a rather rare clone, based on the fact that its messenger RNA was found only in the melanoma cells and not in the other tissues examined like brain, thymus, parathyroid gland, parotid gland, salivary gland, adrenal gland, testis, liver, lung, heart, spleen, skeletal muscle, intestine and colon, cf. Example 1.

Transmembrane segments of the above-mentioned polypeptide (corresponding to nucleotides 286-351, 394-465, 517-588, 640-711, 733-804, 898-972 and 997-1068 in SEQ ID NO: 1, respectively) were determined by hydropathy analysis (Kyte et al. 1982).

Glycosylation sites are found at amino acid residues 15 and 29 in SEQ ID NO: 2, possible phosphorylation sites are found at amino acid residues 42-45, 151-154 and 306-308 SEQ ID NO: 2, and a possible palmitylation site is found at amino acid residue 316 in SEQ ID NO: 2.

The abbreviations of the amino acids used herein are the following:

Amino acid	Three-letter abbreviation	One-letter symbol	
Alanine 5 Arginine Asparagine Aspartic acid Cysteine Glutamine 10 Glutamic acid Glycine Histidine Isoleucine Leucine 15 Lysine Methionine Phenylalanine Proline Serine 20 Threonine Tryptophan Tyrosine Valine	Ala Arg Asn Asp Cys Gln Glu Gly His Ile Leu Lys Met Phe Pro Ser Thr Trp Tyr Val	A R N D C Q E G H I L K M F P S T W Y V	

- Each of the nucleotides shown herein is represented by the abbreviations generally used, i.e.
  - A represents deoxyadenine
  - T represents deoxythymidine
  - G represents deoxyguanine
- C represents deoxycytosine

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N represents deoxyinosine

Using the DNA fragment G-8 described below (with the nucleotide sequence shown in SEQ ID NO: 7) as a hybridization probe, another novel DNA fragment has been isolated from a human genomic library. This DNA fragment is in the present application numbered as SEQ ID NO: 15. This fragment constitutes another interesting aspect of the invention as it has been shown to code for a previously unknown polypeptide which is also an MSH receptor and/or an MSH receptor subtype. The polypeptide encoded by the fragment is in the present context 40 numbered as SEQ ID NO: 16. It is believed that the polypeptide is melanotropic hormone receptor such as an  $\alpha\textsc{-MSH}$  recep-

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tor and/or a  $\beta$ -MSH receptor and/or a  $\gamma$ -MSH receptor and/or an ACTH receptor and is interchangeably referred to as MC-2 and MC-2 receptor herein. The MC-2 receptor is in particular known to be located in the central nervous system and also in peripheral organs such as gut, lung, heart, liver, spleen, smooth and skeletal muscle tissues and the immune system.

The novel DNA fragment with the nucleotide sequence SEQ ID NO: 15 comprises 1650 nucleotides and was sequenced as described herein. The nucleotides from 1 to 615 form the 5' untranslated region while the nucleotides 1591 to 1650 form the 3' untranslated region. The coding fragment from nucleotide 616-1590 encodes a polypeptide of 325 amino acids which is shown in SEQ ID NO: 16. The DNA-fragment was isolated from a human genomic library as described in Example 6.

- Thus, an aspect of the invention relates to a DNA fragment having the nucleotide sequence shown in SEQ ID NO: 15 or an analogue or subsequence thereof which
  - 1) has a homology with the DNA sequence shown in SEQ ID NO: 15 of at least 50%, and/or
- 20 2) encodes a polypeptide, the amino acid of which is at least 50% homologous with the amino acid sequence shown in SEQ ID NO: 16, and/or
  - 3) encodes a polypeptide which binds an antibody which is also bound by an MSH receptor, and/or
- 25 4) encodes a polypeptide which is an MSH receptor or which has the same binding capacity as an MSH receptor.

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While one particular aspect of the invention relates to DNA fragments having the nucleotide sequence shown in SEQ ID NO: 1 or in SEQ ID NO: 15 and encoding a polypeptide of the invention, an analogue or subsequence thereof comprising at

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least 15 nucleotides is another important aspect of the invention. The invention relates to the coding part of the described nucleotide sequence shown in SEQ ID NO: 1 or SEQ ID NO: 15 as well as the non-coding part. A DNA fragment which shows at least 55% homology, preferably at least 70%, more preferably at least 80% and most preferably at least 95% sequence homology with a DNA fragment of the same length obtained from the DNA sequence shown in SEQ ID NO: 1 or SEQ ID NO: 15 is also an interesting aspect of the invention as such fragments and subsequences may encode polypeptides capable of acting as epitopes and thus capable of eliciting an antibody response directed thereto. Such antibodies can also bind to a polypeptide constituting an MSH receptor and thereby being important in diagnosis and treatments of MSH receptor related diseases and conditions, as will appear from 15 the following. In addition, such fragments and subsequences may among other utilities be used as probes in the identification of other DNA fragments as will appear from the following. In this respect a fragment and/or subsequence of the non-coding part of the DNA fragments shown in SEQ ID NO: 1 and 15 is equally important as the fragments and/or subsequences of the coding parts of these DNA fragments.

When used in the present context with regard to nucleotide sequences, the term "subsequence" indicates a nucleotide sequence which is derived from a DNA fragment of the invention and which has retained a characteristic nucleotide sequence thereof as evidenced by its conforming to at least one of the criteria 1)-4) above. Typically, the subsequence is a part of a nucleotide sequence shown in SEQ ID NO: 1 or SEQ ID NO: 15, the subsequence being either a consecutive 30 stretch of nucleotides taken from a nucleotide sequence shown in SEQ ID NO: 1 or SEQ ID NO: 15 or being composed of one or more separate nucleotides or nucleotide sequences of a nucleotide sequence shown in SEQ ID NO: 1 and/or SEQ ID NO: 15.

It is important to note that a "characteristic nucleotide 35 sequence" in the present context is meant to indicate a

nucleotide sequence of a DNA fragment of the invention which is identifying the DNA fragment according to one or more of the following criteria:

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- it encodes a peptide with binding properties of an MSH
   receptor, and/or
  - it encodes a peptide which is bound (with high specificity) by an antibody which also binds (with high specificity) to the polypeptide encoded by the original DNA fragment from which it is derived (the binding for instance being assessed as described in example 10), and/or

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- it will be useful as a hybridization probe for identifying the original DNA fragment from which it is derived.

When a compound "X", such as a e.g. receptor, "binds" to a polypeptide, an antibody or another substance "Y" it is in the present application defined as a substantial specific binding of "X" to "Y" as assessed by the ability of "X" to distinguish between "Y" and other substances under physiological conditions (e.g. in a Ringer solution at 37°C or e.g. using the binding buffer and conditions essentially as described in Example 3). It is preferred that "X" binds "Y" with known affinity. Preferably, the dissociation constant (defined as  $K = \frac{\lambda_X \lambda_Y}{\lambda_{XY}}$ , wherein  $\lambda_X$ ,  $\lambda_Y$  and  $\lambda_{XY}$  are the activities

of "X", "Y" and "XY" in the system XY = X+Y) of the complex

"XY" is less than 10 μM, more preferably less than 1 μM, even more preferably less than 100 nM and most preferably the dissociation constant of the complex "XY" is less than 10 nM. Moreover, in a still further meaning, when "X" is said to bind to "Y", the latter which is also bound to a compound "Z" with a known high affinity (it has been established that "Y" binds to "Z" with such high affinity that it can be regarded as a test-tool), the dissociation constant of the complex "XY" formed is not more than 100,000-fold higher than that

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for "YZ", preferably not more than 10,000-fold, more preferably not more than 1,000-fold and most preferably not more than 100-fold higher than that for "YZ".

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One way of determining such binding characteristics is to 5 employ the method of example 3, wherein a panel of melanotropic hormones are tested with regard to their ability to inhibit binding of  $^{125}I-NDP-MSH$  to a suspected MSH receptor.

According to the above, a degree of homology of as little as 30% between a DNA fragment and one of the DNA fragments shown in SEQ ID NO: 1 or 15 or a subsequence thereof may in some 10 instances ensure that a characteristic DNA sequence is retained in the first DNA fragment. Thus, the invention also relates to DNA fragments which have retained a characteristic DNA sequence of the DNA sequences in SEQ ID NO: 1 and 15, said characteristic DNA sequences having as little as 30% homology with any of the sequences shown in SEQ ID NO: 1 and 15. This will most likely be the case when the characteristic sequence is a part of a functional important part of the polypeptide and therefore has little resemblance with other 20 polypeptides. However, it is preferred that the degree of homology is at least 40% when the characteristic sequence codes a less functionally important part of the molecule. The degree of homology may in some instances be so high as 95%; this might be the case when the characteristic DNA fragment encodes a intramembraneous part of the polypeptide, wherein the amino acid sequence might be much like other sequences of transmembraneous polypeptides.

In the present specification and claims, the term "subsequence" thus designates a nucleotide sequence which pre-30 ferably has a size of at least 15 nucleotides, more preferably at least 18 nucleotides, still more preferably at least 21 nucleotides, even more preferably at least 27 nucleotides and most preferably at least 51 nucleotides. It is well known that small fragments are useful as epitopes, DNA-probes for hybridization with DNA or RNA, in PCR techniques as is de-

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scribed herein, or useful in that they encode peptides comprising epitopes capable of eliciting the production of antibodies.

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The term "analogue" with regard to the DNA fragments of the invention is intended to indicate a nucleotide sequence which encodes a polypeptide identical or substantially identical to a polypeptide encoded by a DNA fragment of the invention shown in SEQ ID NO: 1 or 15.

It is well known that the same amino acid may be encoded by various codons, the codon usage being related, inter alia, to the preference of the organisms in question expressing the nucleotide sequence. Thus, one or more nucleotides or codons of a DNA fragment of the invention may be exchanged by others which, when expressed, result in a polypeptide identical or substantially identical to the polypeptide encoded by the DNA fragment in question.

Thus, a DNA fragment encoding a polypeptide comprising the amino acids 1-317 of SEQ ID NO: 2, or a DNA fragment encoding a polypeptide comprising the amino acids 1-325 SEQ ID NO: 16 are very important embodiments of the invention.

Also, the term "analogue" is used in the present context to indicate a DNA fragment or a DNA sequence of a similar nucleotide composition or sequence as the DNA sequence encoding the amino acid sequence constituting an MSH receptor, allowing for minor variations which do not have an adverse effect on the ligand binding properties and/or biological function and/or immunogenicity as compared to the MSH receptor, or which give interesting and useful novel binding properties or biological functions and immunogenicities etc.

30 of the analogue. The analogous DNA fragment or DNA sequence may be derived from an animal or a human or may be partially or completely of synthetic origin as described above. The analogue may also be derived through the use of recombinant DNA techniques.

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Furthermore, the terms "analogue" and "subsequence" are intended to allow for variations in the sequence such as substitution, insertion (including introns), addition, deletion and rearrangement of one or more nucleotides, which 5 variations do not have any substantial effect on the polypeptide encoded by a DNA fragment or a subsequence thereof. The term "substitution" is intended to mean the replacement of one or more nucleotides in the full nucleotide sequence with one or more different nucleotides, "addition" is under-10 stood to mean the addition of one or more nucleotides at either end of the full nucleotide sequence, "insertion" is intended to mean the introduction of one or more nucleotides within the full nucleotide sequence, "deletion" is intended to indicate that one or more nucleotides have been deleted from the full nucleotide sequence whether at either end of the sequence or at any suitable point within it, and "rearrangement" is intended to mean that two or more nucleotide residues have been exchanged with each other.

When using the term "any substantial effect on the 20 polypeptide" is understood that the DNA fragment encodes a polypeptide which has retained its antigenicity and/or MSH binding properties compared to the MSH receptor polypeptide encoded by the DNA fragment from which the analogue/subsequence is derived.

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The terms "fragment", "sequence", "subsequence" and "analogue", as used in the present specification and claims with respect to fragments, sequences, subsequences and analogues according to the invention should of course be understood as not comprising these phenomena in their natural environment, 30 but rather, e.g., in isolated, purified, in vitro or recombinant form.

The terms "homology" and "homologous" are, with respect to DNA fragments, intended to mean a homology between the nucleotides in question between which the homology is to be 35 established, in the match with respect to identity and posi-

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tion of the nucleotides of the DNA fragments. With respect to polypeptides and fragments thereof described herein, the terms are intended to mean a homology between the amino acids in question between which the homology is to be established, in the match with respect to identity and position of the amino acids of the polypeptides.

"Binding capacity of an MSH receptor" is in this context meant as the binding properties of an MSH receptor assessed by a test wherein the binding between the receptor and various possible ligands is determined with respect to their ability of inhibiting binding of <sup>125</sup>I-NDP-MSH (<sup>125</sup>I-(Nle<sup>4</sup>,D-Phe<sup>7</sup>)-MSH) as described herein.

"Same binding capacity as an MSH receptor" is defined herein as a binding profile which shows that a substance binds <sup>125</sup>I-NDP-MSH with a higher affinity than other compounds although the substance may be unable to elicit the effects exerted by the binding of MSH to an MSH receptor.

When reference is being made to "an analogue of MSH" or "an analogue of a melanotropic hormone" it is intended to mean a substance which shows binding capacity for an MSH receptor as defined above. Thus, examples of analogues of MSH and analogues of melanotropic hormones are  $\alpha$ -MSH,  $\beta$ -MSH,  $\gamma$ -MSH and NDP-MSH.

The term "melanotropic hormone" is intended to refer both to a natural peptide being derived from proopiomelanocortin (POMC), the natural peptide typically having a biological activity of that of MSH or ACTH, and a synthetic peptide, the synthetic peptide having the ability to induce at least one of the biological effects which may be induced by the natural melanotropic hormones. Examples of melanotropic hormones are  $\alpha$ -MSH,  $\beta$ -MSH,  $\gamma$ -MSH, ACTH and NDP-MSH.

The term "melanotropic hormone receptor" is intended to mean a receptor which can be activated by a melanotropic hormone

so as to induce a second messenger response (or any other typical receptor response) or a biological effect generally being referred to as a melanotropic hormone response. Stimulation of melanin formation in melanocytes and the stimulation of corticosteroid synthesis in the adrenal gland by the melanotropic hormones are typical examples of melanotropic hormone responses. Examples of melanotropic hormone receptors are the MSH receptor, the ACTH receptor and the MC-2 receptor.

- 10 The present invention is based on the construction of the primers shown as SEQ ID NO: 3 and SEQ ID NO: 4 and the analogues thereof which are defined below. As appears from the above, great interest and many efforts have been exerted in order to examine the function of the MSH receptor and thus, there has been obvious interest in isolating the DNA encoding the MSH receptor. The very careful work performed by the inventors of the present invention when designing these primers such as described in details in Example—1 rendered the present invention possible.
- Thus, the DNA fragments of the invention used as primers 20 constitute another interesting aspect of the invention and have various important utilities such as detection and isolation of other DNA fragments encoding polypeptides having similar functions and/or binding capacity as an MSH receptor. In particular, the primers can be used in the detection of 25 other G-protein coupled or binding receptors. The invention therefore also relates to a DNA fragment having the nucleotide sequence SEQ ID NO: 3 (from segment 3) or analogues thereof, wherein the nucleotides 13 and/or 15 and/or 23 optionally are substituted by C and to a DNA fragment having the nucleotide sequence SEQ ID NO: 4 (from segment 6) or analogues thereof wherein the nucleotides 19 and/or 29 and/or 32 optionally are substituted by C, and wherein the nucleotides 20 and/or 31 are optionally substituted by G.

In order to examine a DNA fragment of the invention or an analogue or subsequence thereof or an RNA fragment transcribed therefrom, such as to examine the relatedness to other foreign DNA fragments, hybridization is a useful method.

5 Hybridization may be performed as follows: A DNA fragment or an analogue or a subsequence thereof of the invention is

an analogue or a subsequence thereof of the invention is labelled with any of the labelling principles available (radioactive system, colour reaction system, light based system, or variations of these) so as to constitute a probe.

The foreign DNA/RNA to be examined is coupled to a matrix.

The matrix is subjected to a suitable treatment so as to couple the DNA/RNA to the matrix. The matrix is exposed to a prehybridization solution of a composition, at a temperature and for a period of time suited to the matrix and the foreign

DNA/RNA in question. The matrix is then placed in a hybridization solution containing labelled denatured DNA probe.

Hybridization is carried out at a suitable temperature and the period of time. The matrix is then washed with a solution of a composition, at a temperature and for a period of time

suited to the matrix and the foreign DNA/RNA in question. The matrix is then subjected to a suitable detection system based on the nature of the label in the DNA probe. The results are then analyzed. Any hybridization of the foreign DNA/RNA and the DNA probe is an indication of similarity of the two

species, and may be used to examine whether the foreign DNA/RNA is a part of the invention. In the above hybridization procedure a RNA probe corresponding to a polypeptide or an analogue or a subsequence thereof of the invention can also be used in place of a DNA probe. Another approach of

determining similarity between DNA sequences is by determining the nucleotide sequence of the DNA fragment to be compared with a DNA fragment or an analogue or subsequence thereof of the invention by conventional DNA sequencing analysis, and comparing the degree of homology with the DNA fragment or an analogue or subsequence thereof of the inventional

fragment or an analogue or subsequence thereof of the invention.

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Polymerase chain reaction (PCR) primers can be synthesized based on the nucleotide sequence of the cloned MSH receptor, or on the basis of other known similar sequences. These primers can then be used to amplify the whole or a part of an MSH receptor sequence or the sequences of its analogues. Primers as shown in SEQ ID NO: 3 and SEQ ID NO: 4 which constitute part of the invention may be used in this aspect. Polymerase chain reaction enzyme, a type of heat stable DNA polymerase, generally incorporates wrong nucleotides at a frequency of 1 in 10000 (Tindall et al. 1988) during amplification. Because of the iterative nature of the amplification this frequently attributes a new altered sequence to the amplified MSH receptor.

The DNA fragment described above and constituting an impor-15 tant aspect of the invention may be obtained directly from the genomic DNA or by isolating mRNA and converting it into the corresponding DNA sequence by using reverse transcriptase, thereby producing a cDNA. When obtaining the DNA fragment from genomic DNA, it is derived directly by screening for 20 genomic sequences such as is described in Example 1. It can be accomplished by hybridization to a DNA probe designed on the basis of knowledge of an MSH receptor sequence, or the sequence information obtained by amino acid sequencing of the purified MSH receptor. When the DNA is of complementary DNA (cDNA) origin, it may be obtained by preparing a cDNA library 25 with mRNA from cells containing MSH receptor or parts thereof. Hybridization can be accomplished by a DNA probe designed on the basis of knowledge of an MSH receptor sequence, or the sequence information obtained by amino acid sequencing of the purified MSH receptor. 30

The DNA fragments of the invention or analogues or subsequences thereof can also be obtained using other methods (Wright et al. 1992) like expression cloning in cell line (Xie et al. 1992) or the expression cloning in the oocyte (Julius et al. 1988; Masu et al. 1987).

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A DNA fragment of the invention or an analogue or subsequence thereof can be obtained from other animals, such as mammals. The DNA obtained in this way could be exactly similar to the one shown in SEQ ID NO: 1 or SEQ ID NO: 15 or could have differences in structure attributed to well known interspecies variations.

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A DNA fragment of the invention or an analogue or subsequence thereof can be replicated by fusing it with a vector and inserting the complex into a suitable microorganism or a mammalian cell line. Alternatively, the DNA fragment can be manufactured using chemical synthesis.

As mentioned above the polypeptides encoded by the DNA fragments of the invention shown in SEQ ID NO: 1 and 15 have been shown to contain coding regions encoding an MSH receptor and an MSH receptor/MSH receptor subtype, respectively.

Thus, in another particular important aspect, the invention relates to a polypeptide having the amino acid sequence shown in SEQ ID NO: 2 or an analogue or subsequence thereof which

- 1) is an MSH receptor or which is capable of binding to MSH or an analogue thereof, and/or
  - 2) is encoded by a DNA fragment which is at least 50% homologous with the DNA fragment shown in SEQ ID NO: 1, and/or
  - 3) binds an antibody which is also bound by an MSH receptor.
- 25 Another most important aspect of the invention thus relates to a polypeptide having the amino acid sequence shown in SEQ ID NO: 16 or an analogue or subsequence thereof which
  - 1) is an MSH receptor or which is capable of binding to MSH or an analogue thereof, and/or

2) is encoded by a DNA fragment which is at least 50% homologous with the DNA fragment shown in SEQ ID NO: 1, and/or

3) binds an antibody which is also bound by an MSH receptor.

being capable of binding MSH and by the term "analogue thereof" is meant any polypeptide having the same binding capacity
as an MSH receptor in that the polypeptide is capable of
binding MSH. Thus, included is also a polypeptide from different sources, such as different animals, such as mammals, in
particular a human, which vary for example in the carbohydrate part, or the phosphorylation and/or in tissue distribution. In this context the term MSH receptor also refers to
both the above-mentioned polypeptides of the invention.

The term analogue also includes polypeptides being capable of binding antibodies which also bind to an MSH receptor. Such analogues may be capable of eliciting or stimulating an immune response which is also directed against the MSH receptor or which can also be elicited by the MSH receptor. These and other analogues are encoded by a DNA fragment or analogue or subsequence thereof of the invention which with respect to analogues have been defined above. The analogues may in a particular aspect be of synthetic origin as discussed herein.

The term "analogue" with regard to a polypeptide is also used in the present context to indicate a protein or polypeptide of a similar amino acid composition or sequence as the characteristic amino acid sequence shown in SEQ ID NO: 2 or SEQ ID NO: 16 (or another polypeptide of the invention), allowing for minor variations which do not have an adverse effect on the ligand binding properties and/or biological function and/or immunogenicity, or which may give interesting and useful novel binding properties or biological functions and immunogenicities etc. of the analogue. The analogous polypeptide or protein may be derived from an animal or a human

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or may be partially or completely of synthetic origin. The analogue may also be derived through the use of recombinant DNA techniques.

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It is being widely recognized that the same or similar gene may be present in two or several copies in the genome of the same animal. Because gene mutations will always tend to induce divergence of a DNA sequence the structure of the protein coded by the same and/or similar genes will tend to diverge during evolution. Thus, in the present context it is 10 obvious that the MSH receptor, because of the existence of the DNA according to SEQ ID NO: 1 and SEQ ID NO: 15 in fact exist in at least two copies in the genome. Thus SEQ ID NO: 1 and 15 are both coding for proteins which have MSH receptor properties, albeit the binding properties of the two proteins for melanotropic hormones differ. It is therefore predicted that even more copies of the MSH receptor are present in the genome of an animal, for instance in homo sapiens. Because of the similarity between the amino acid composition of such proteins and the MSH receptors described herein, theses 20 proteins are in the present application being regarded as being analogues of the MSH receptor. Thus such proteins are also part of the invention.

In the present context the term "characteristic amino acid sequence derived from an MSH receptor" is intended to mean an amino acid sequence which comprises amino acids constituting a substantially consecutive stretch (in terms of linear or spatial conformation) of the polypeptide shown in SEQ ID NO: 2 or the amino acid sequence shown in SEQ ID NO: 16 and encoding an MSH receptor. Such secondary or tertiary conformation may have interesting and useful properties and may 30 constitute epitopes.

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In the present context, the term "epitope" refers to any polypeptide of the invention or an analogue thereof capable of stimulating or interacting with immunocompetent cells and capable of stimulating the production of antibodies which

also bind to a polypeptide constituting an MSH receptor. Especially epitopes showing desirable properties with regard to diagnosis and therapy constitute important aspects of the present invention.

In the present context, the term "epitope" also refers to any polypeptide of the invention or a characteristic amino acid sequence or an analogue thereof capable of interacting or binding existing or novel substances which are also bound by a polypeptide constituting an MSH receptor. The said substances can be organic molecules, small peptides or large polypeptides or derivatives of any of the above. Such an approach can find use in the drug screening programme.

The term "receptor subtype" is intended to mean a receptor which is capable of binding the same ligand and/or ligands as another receptor, albeit the affinities of the ligands for the receptors may be different for the compared receptors.

Thus, when referring to MC-2 as an "MSH receptor subtype" it is indicated that MC-2 is an MSH receptor, but that the pattern of binding to various ligands/substances is different from that of another MSH receptor, such as e.g. the MSH receptor with the amino acid sequence shown in SEQ ID NO: 2.

The term "subsequence" with regard to a polypeptide designates a polypeptide sequence which comprises a part of the polypeptide sequence shown in SEQ ID NO: 2 or SEQ ID NO: 16 or other polypeptide sequences of the invention which may optionally have retained its capability of binding MSH.

Included are also polypeptide subsequences which have been analogized by modifications as explained herein. Polypeptides constituting interesting epitopes or encoded by a nucleotide subsequence of the invention as defined above are also included.

"A derivative of an MSH receptor" is meant to indicate both an analogue, subsequence or subtype of an MSH receptors.

In a most important aspect, the invention relates to a polypeptide encoded by the DNA fragment shown in SEQ ID NO: 1, preferably the polypeptide shown in SEQ ID NO: 2, and to a polypeptide encoded by the DNA fragment with the nucleotide sequence shown in SEQ ID NO: 15, preferably the polypeptide shown in SEQ ID NO: 16.

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The polypeptides of the invention also comprises polypeptides which show a degree of homology of at least 55%, preferably at least 70%, more preferably at least 80% and most preferably 10 ly at least 95% homology to a polypeptide of the same length which has an amino acid sequence which is a part of the sequences shown in SEQ ID NO: 2 and SEQ ID NO: 16.

The invention also relates to a characteristic amino acid sequence being a subsequence comprising from at least 5 amino acids to 316 acids of SEQ ID NO: 2 and to a subsequence comprising from 5 to 324 amino acids of SEQ ID NO: 16, and any analogue to such polypeptides. Preferably the subsequence comprises at least 7 amino acids, more preferably at least 10 amino acids, even more preferably at least 15 amino acids and most preferably at least 30 amino acids. The polypeptide may be coupled to any other moiety.

The present invention also relates to a substantially pure polypeptide which has the same binding capacity as an MSH receptor or which is recognized by an antibody raised against or reactive with a polypeptide of the invention. Furthermore the invention relates to any polypeptide of the invention in substantially pure form.

furthermore, the invention relates to a polypeptide as defined herein which is glycosylated or which is linked to a carbohydrate or lipid moiety. Also a polypeptide containing a palmitoyl anchor or a part thereof constitutes an interesting aspect as well as any polypeptide of the invention in lipid soluble form which may, in one interesting aspect of the invention as described herein, be used in the treatment of an

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animal, in particular a human, having impaired function of the receptor.

The lipid soluble form of polypeptide of the invention may be a form comprising components such as liposomes, micelles and phospholipid so as to allow the polypeptide to be incorporated in the cell membrane of the recipient. It is important that the lipid soluble form is a form which ensures the stability of the polypeptide and preferably in a form which is pharmaceutically acceptable so as to allow the administration of the lipid soluble form to an animal, in particular a human. The lipid soluble form may also be in a form comprising components such a detergent, oil, such as mineral oil or vegetable oil or water, and which may be a suspension of one or more of the above mentioned components.

In the present context, the term "substantially pure" is understood to mean that the polypeptide in question is substantially free from other components, e.g. other polypeptides or carbohydrates, which may result from the production and/or recovery of the polypeptide or otherwise be found together with the polypeptide. The high purity of a polypeptide of the invention is advantageous when the polypeptide is to be used for, e.g., the production of antibodies. Also due to its high purity, the substantially pure polypeptide may be used in a lower amount than a polypeptide of a conventional lower purity for most purposes. The purification of a polypeptide of the invention may be performed by methods known to a person skilled in the art.

The polypeptides of the invention having the amino acid sequences shown in SEQ ID NO: 2 and 16 and which are MSH receptors bear similarity with other G-protein coupled receptors. They have the most common feature of passing through the cell membrane 7 times, like all other G-protein coupled receptors. Based on the observations of homology between the transmembrane segments of different G-protein coupled receptors, it has been hypothesized that the extra-

cellular loops and the transmembrane segments are involved in the ligand binding, such as e.g. the binding of MSH or an analogue of MSH or of a synthetic organic molecule serving as ligand for the receptor. The intracellular loops have been assigned the role of coupling to the G-proteins and possible involvement in other intracellular activities. In the present context the above mentioned extra and intracellular loops as

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well the transmembrane segments that are intended to be involved in the binding of the ligand and/or the coupling of G-proteins are one or two or several or all of the mentioned segments and loops of the receptor.

Intense efforts to solve the 3-dimensional (3D) structure of G-protein coupled receptors and some related proteins (e.g. bacteriorhodopsin and opsins) are ongoing in several laboratories world wide. It is expected that once the 3D structure of one of these proteins is solved, the 3D structure of other G-protein coupled receptors will become easily solvable using computational methods, provided that their primary amino acid structure is known. This is due to the fact that all G-protein coupled receptors are likely to show similar gross 3D structure (Sankara-Ramakrishnan & Vishveshwara 1989; Findlay & Eliopoulos 1990; Hibert M.F. et al. 1991).

Thus, in certain embodiments of the invention are considered especially important amino acids 1-39, 100-116, 182-188 and 269-276 which are considered to constitute the extracellular loops of the MSH receptor. These regions will be of particular importance as epitope targets for antibodies intended for clinical use in e.g. targeted drug delivery. or for drug design.

In other embodiments of the invention are considered especially important the amino acids 63-75, 142-157, 213-243 and 301-317 which are considered to constitute the intracellular loops of the MSH receptor. These regions will be particularly important in the elucidation of the mechanisms for the coupling of the MSH receptor to G-proteins. These regions may

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serve as targets for drugs aiming for the modulation of the interaction of the receptor with G-proteins. It may also in particular be desired to alter the amino acid sequences, by e.g. deletions, site directed mutations, insertions of extra 5 amino acids, or combinations thereof, to generate novel MSH receptor analogues showing altered properties. Such altered receptors may be desired to further the understanding of the molecular mechanisms in the coupling of the receptor with G-proteins. Moreover, yet another aspect pertaining to this 10 particular part of the invention is the DNA sequences coding for the intracellular segments, in particular base pairs 636-726, but also the other segments, as these regions are considered to be less homologous with other G-protein coupled receptor coding fragments. Thus, such regions of the sequence 15 may serve to generate DNA probes which in hybridization studies, as is described in detail below, are selective for the MSH receptor DNA or mRNA.

In yet other embodiments of the invention are considered particularly important the amino acids 40-62, 76-99, 117-140, 20 158-181, 189-212, 244-268 and 277-300 which are considered to constitute the transmembrane segments of the MSH receptor. It may here be desired to alter one or several specific amino acids to generate MSH receptor analogues showing altered properties. Such altered receptors may be desired to further the understanding of the molecular mechanisms in the binding of MSH and MSH analogues to the receptor. Moreover, yet another aspect pertaining to this particular part of the invention is the DNA sequences coding for the transmembrane segments as these regions are considered to be highly homologous with other G-protein coupled receptor coding fragments 30 which are natural variants of the MSH receptor. Such receptor coding fragments may exist in other species which code for species variants of the MSH receptor. Such receptor coding fragments may also exist in humans as well as animals which encode homologous receptors which are subtypes of the MSH 35 receptor or which are closely related receptor types possibly belonging to the same class of melanotropic hormone receptor

family. By using homology screening methods utilizing DNA sequences derived from the transmembrane segments it may be possible to obtain the DNA sequences of these homologous receptor coding fragments.

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Because of the difference in sequence of SEQ ID NO: 16 from SEQ ID NO: 2 the following fragments of the MC-2 receptor are considered especially important: Amino acids 1-37, 98-114, 180-186 and 266-273 of SEQ ID NO: 16 which are considered to constitute the extracellular loops of the MC-2 receptor.

These regions will be of particular importance as targets for antibodies intended for clinical use in e.g. targeted drug

delivery or for drug design.

In other embodiments of the invention are considered especially important the amino acids 62-73, 139-155, 212-239 and 298-325 of SEQ ID NO: 16 which are considered to constitute 15 the intracellular loops of the MC-2 receptor. These regions will be particularly important in the elucidation of the mechanisms for the coupling of the MC-2 receptor to G-proteins. These regions may serve as targets for drugs aiming for 20 the modulation of the interaction of the receptor with G-proteins. It may also in particular be desired to alter the amino acid sequences, by e.g. deletions, site directed mutations, insertions of extra amino acids, or combinations thereof, to generate novel MC-2 receptor analogues showing altered properties. Such altered receptors may be desired to 25 further the understanding of the molecular mechanisms in the coupling of the receptor with G-proteins. Moreover, yet another aspect pertaining to this particular part of the invention is the DNA sequences coding for the intracellular segments, in particular base pairs 633-717 of SEQ ID NO: 15, 30 but also the other segments, as these regions are considered to be less homologous with other G-protein coupled receptor coding fragments. Thus, such regions of the sequence may serve to generate DNA probes which in hybridization studies, as is described in detail below, are selective for the MC-2 receptor DNA or mRNA.

In yet other embodiments of the invention are considered particularly important the amino acids 38-61, 74-97, 115-138, 156-179,187-211, 240-265 and 274-297 of SEQ ID NO: 16 which are considered to constitute the transmembrane segments of 5 the MC-2 receptor. It may here be desired to alter one or several specific amino acids to generate MC-2 receptor analogues showing altered properties. Such altered receptors may be desired to further the understanding of the molecular mechanisms in the binding of MSH and MSH analogues to the 10 receptor. Moreover, yet another aspect pertaining to this particular part of the invention is the DNA sequences coding for the transmembrane segments as these regions are considered to be highly homologous with other G-protein coupled receptor coding fragments which are natural variants of the 15 MC-2 receptor. Such receptor coding fragments may exist in other species which code for species variants of the MC-2 receptor. Such receptor coding fragments may also exist in humans as well as animals which encode homologous receptors which are subtypes of the MC-2 receptor or which are closely 20 related receptor types possibly belonging to the same class of melanotropic hormone receptor family. By using homology screening methods utilizing DNA-sequences derived from the transmembrane segments it may be possible to obtain the DNA sequences of these homologous receptor coding fragments.

The above mentioned specific amino acid sequences are prominent examples of subsequences according to the invention. It is to be understood that the other important subsequences according to the invention are subsequences which are modifications of the above mentioned subsequences in that, and with respect to particular subsequences of which they are modifications, they fulfil any one of the criteria 1)-3) for the polypeptide as stated above. Also included in this aspect of the invention is a DNA fragment encoding any such amino acid sequence.

Using the primers of the invention, three DNA fragments have been isolated and sequenced. These DNA fragments termed G-6

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shown in SEQ ID NO: 5, G-8, shown in SEQ ID NO: 7 and G-10, shown in SEQ ID NO: 9, share homologies with the DNA sequence shown in SEQ ID NO: 1.

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Thus, the present invention also relates to a DNA fragment or a subsequence or an analogue thereof which shows a homology with any of the nucleotide sequences shown in SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9 of at least 40%, or which can be isolated by using the nucleotide sequence shown in SEQ ID NO: 13 and/or SEQ ID NO: 14 as a primer, or which has any of the nucleotide sequences shown in SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9. The homology may in some aspects be at least 50%, preferably at least 55%, more preferably at least 70%, even more preferably at least 80% and most preferably at least 95% with any of the DNA sequences shown in SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9, respectively.

Polypeptide sequences or subsequences or analogues thereof which show a homology of at least 40% with any of the polypeptides shown in SEQ ID NO: 6, 8 or 10 encoded for by the DNA sequences shown in SEQ ID NO: 5, SEQ ID NO: 7 and SEQ ID NO: 9, respectively, constitute yet another embodiment of the invention. The homology may in some aspects be at least 50%, preferably at least 55%, more preferably at least 70%, even more preferably at least 80% and most preferably at least 95% with any of the polypeptides shown in SEQ ID NO: 6, SEQ ID NO: 8 and SEQ ID NO: 10, respectively.

Any of the DNA fragments of the invention may in a particular embodiment of the invention be used for the preparation of a DNA probe which may be labelled or unlabelled and which is used to isolate full length coding fragments and/or to detect or quantitate RNA. The information of any of these DNA fragments may also be used to make PCR primers and for making a polypeptide.

As mentioned above and described in the examples, the G-8 fragment has been used to isolate a new MSH receptor subtype,

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MC-2. It is therefore contemplated to perform similar experiments using the two other fragments G-6 and G-10.

The DNA fragments G-6 and G-10 represent two potential G-protein coupled receptors, and can be used to isolate full length coding fragments for them using for example the same methodology as described in Example 1.

In particular, because of substantial homology of G-6 and G-10 with the primers shown in SEQ ID NO: 3 and SEQ ID NO: 4 of the invention, it is considered that one or both of G-6 and G-10 represent coding fragments encoding receptors for peptides sharing similar origin as the MSH receptor; that is the POMC (proopiomelanocortin) receptors, such as e.g. the ACTH receptor, the met-enkephalin receptor and the  $\beta$ -endorphin receptor. In particular, it is considered that one or both of G-6 and G-10 are the  $\mu$ ,  $\delta$  and  $\sigma$  opioid receptors because of their dissimilarities to the  $\kappa$  opioid receptor.

However, G-6 and G-10 each has unique sequence characteristics starting in the intracellular segment 5, that is the amino acids "L-Y-V/I-H-M", and it may be contemplated that one or both of G-6 and G-10 represent parts of DNA sequences encoding receptors for a novel class of peptide hormone receptors; e.g. the corticotropin releasing hormone receptor, the growth hormone releasing hormone receptor, the gonadotropin releasing hormone receptor, the thyrotropin releasing hormone receptor, the luteinizing hormone releasing hormone receptor, the follicle stimulating hormone releasing hormone receptor, the chorionic gonadotropin hormone receptor and/or the glucagon receptor.

However, G-6 and G-10 also has short extracellular loops, indicating that the receptors encoded by these fragments constitute the small peptide hormone type of G-protein coupled receptors. Thus, one or both of G-6 and G-10 may represent the neuropeptide Y receptor, the tumour necrosis factor receptor, the colony stimulating factor receptor, the

interleukin 1 receptor, the neurotensin receptor, the atrial natriuretic factor receptor, the kallidin receptor, the bulbogastrin receptor, the motilin receptor, the pancreatic polypeptide receptor, the olfactory receptor subtype, the 5 spermatozoon feromone receptor subtype, the insulin like growth factor receptor, the taste receptor subtype, the gustuducin coupled receptor subtype, the inhibin receptor subtype and/or the kyotropin receptor.

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The dissimilarity of the G-6 and G-10 with other known 10 nucleotide fragments coding for G-protein coupled receptors, the latter of which include a substantial amount of well characterized receptors for amines, may indicate that one or both of G-6 and G-10 by contrast encode genes for lipids, e.g. more specifically receptors for prostanoids, that is, 15 more specifically the prostaglandin  $E_1$  receptor, the prostaglandin  $E_2$  receptor, the prostaglandin  $F_{2a}$  or receptors for prostacyclins, e.g. the PGI2 receptor, and/or receptors for leukotrienes, e.g. the leukotriene D4 receptor and/or the leukotriene C4 receptor and/or a receptor for a non-lipid, 20 e.g. the phosphatidic acid receptor and/or the thromboxane A2 receptor and/or the platelet activating factor receptor.

Due to general characteristics of G-6 and G-10 being G-protein coupled receptors, it is considered that one or both of G-6 and G-10 by contrast encode genes for subtype of the substance P receptor, substance K receptor, endothelin receptor, angiotensin receptor, chemoattractant peptide receptor, bombesin receptor, oxytocin receptor, vasopressin receptor, antidiuretic hormone receptor, gastrin receptor, cholecystokinin receptor, canabinoid receptor, follicle stimulating hormone receptor, luteinizing hormone receptor, 30 growth hormone receptor, thyrotropin receptor, calcitonin receptor, calcitonin gene related peptide receptor and/or parathyroid hormone receptor.

Based on the knowledge of the DNA fragments of the invention (or the knowledge of the analogues or RNA fragments) the 35

described DNA fragments of the invention can be produced containing one or more modified nucleotides to improve resistance against nucleases or to improve transport across cell membranes.

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5 Suitable polypeptides can be produced using recombinant DNA technology. More specifically, the polypeptides may be produced by a method which comprises cultivating or breeding an organism carrying a DNA fragment or an analogue or a subsequence thereof of the invention under conditions leading to expression of said DNA fragment, and subsequently recovering the expressed polypeptide from the said organism.

The organism which can be used for the production of such a polypeptide may be a higher organism e.g. an animal, or a lower organism e.g. a microorganism. Irrespective of the type of organism used, a DNA fragment of the invention or an analogue or a subsequence thereof (described above) should be introduced in the organism either directly or with the help of a suitable vector. Alternatively, the polypeptides may be produced in the mammalian cell lines by introducing a DNA fragment of the invention either directly or with the help of an expression vector.

The DNA fragments or analogues or subsequences thereof of the invention can also be cloned in a suitable stable expression vector and then put into a suitable cell line. The cells expressing the desired polypeptides are then selected using the conditions suitable for the vector and the cell line used. The selected cells are then grown further and form a very important and continuous source of the desired polypeptides.

A polypeptide of the invention can also be made by in vitro translation of the RNA complementary to a DNA fragment of the invention. This can be achieved for the whole molecule, or a part or parts of the molecule, in free form or in fusion with one or several proteins. The methods which can be used are

described (Sambrook et al. 1989; Spirin et al. 1988). The polypeptides of the invention can also be expressed in vitro

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as functional proteins in the fused or the unfused form (Zozulya et al. 1990).

5 In line with the above, the invention relates to a vector containing a recombinant DNA insert coding for an MSH receptor polypeptide of the invention or a fusion polypeptide as defined herein. In one particular important embodiment, a DNA fragment or an analogue or subsequence thereof of the invention or a fusion DNA fragment of the invention as 10 defined herein may be carried by a replicable expression vector which is capable of replicating in a host organism or a cell line.

The vector may in particular be a plasmid, phage, cosmid, 15 mini-chromosome or virus. In an interesting embodiment of the invention, the vector may be a vector which, when introduced in a host cell, is integrated in the host cell genome.

Included as an important aspect of the invention is also an organism which carries and is capable of expressing a DNA 20 fragment of the invention. Such a plasmid vector has been constructed and is designated pE-11D herein. This vector constitutes yet another aspect of the invention.

Also, the invention relates to an organism which carries and is capable of replicating a DNA fragment of the invention and 25 also such a plasmid vector designated pB-11D has been constructed and constitutes a part of the invention. This vector was deposited on 24 August 1992 under the number DSM 7214 in the Deutsche Sammlung von Mikroorganismen under the terms and conditions of the Budapest Treaty.

30 Further, yet another plasmid vector, designated pB-MC-2, has been constructed, which is capable of replicating a DNA of the present invention. pB-MC-2 thus constitutes yet a very

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important aspect of the invention. The construction of pB-MC-2 is described in detail in Example 6.

Moreover, still yet another plasmid vector, designated pE-MC-2, which is also a very important embodiment of the invention, has been constructed. The pE-MC-2 plasmid vector is capable of expressing the MC-2 receptor of the present invention. The construction of pE-MC-2 is detailed in Example 7 and the use of pE-MC-2 for the expression of the MC-2 receptor is exemplified in Example 8. This vector has been deposited on 9 August 1993 under the number DSM 8440 in the Deutsche Sammlung von Mikroorganismen under the terms and conditions of the Budapest Treaty.

Organisms which may be used in this aspect of the invention of producing the peptides of the invention comprise a cell which is a microorganism such as a bacterium, a yeast, a protozoan, or which is derived from a multicellular organism such as a fungus, an insect, a plant, a mammal or it may be a cell line. If the organism is a bacterium, it is preferred that the bacterium is of the genus Bacillus, e.g. B. subti
lis, Escherichia, e.g. E. coli, or Salmonella.

If a higher organism is used, transgenic techniques may be employed for the production of the polypeptides. Examples of suitable animals are sheep, cattle, pigs etc. A DNA fragment encoding a polypeptide of the invention is expressed together with a polypeptide which is inherently expressed by the animal, e.g. a milk protein or the like. The resulting fusion protein may then be subjected to post-translational modifications so as to obtain a polypeptide of the invention.

In another aspect of the invention a MSH receptor may be obtained from a suitable cell type found to naturally express MSH receptor from DNA encoding an MSH receptor. Such cells may be e.g. a melanoma cell line, as is shown in Example 3 for WM 266-4 cells, or they may be obtained from any tissue containing cells expressing a DNA fragment of the invention.

A stable cell line capable of producing a polypeptide of the invention having MSH binding properties, has been established. The cell lines of COS-7 cells constitute other important aspects of the invention. The cell line harbours the DNA fragment with the nucleotide sequence SEQ ID NO: 1 and steadily produces polypeptides having binding properties substantially identical to the binding properties described below. The production of this stable cell line is described in detail in example 5.

Thus, the invention also relates to a stable cell line producing a polypeptide of the invention which optionally binds NDP-MSH with high affinity; the establishment of such a cell line may be performed according to the technique described in Example 5, or to any other method known to the person skilled in the art.

In one particular aspect of the invention, a DNA fragment of the invention may comprise one or more second nucleotide sequence(s) encoding one or more polypeptide(s) different from or identical to a polypeptide of the invention fused in 20 frame to a DNA fragment of the invention or an analogue thereof encoding a polypeptide or an analogue or subsequence thereof of the invention with the purpose of producing a fused polypeptide which polypeptide constitutes yet another interesting aspect of the invention. When using recombinant DNA technology the fused DNA sequences may be inserted into a suitable vector or genome. Alternatively, one of the nucleotide sequences is inserted into the vector or genome already containing the other nucleotide sequence(s). A fusion polypeptide can also be made by inserting the nucleotide sequen-30 ces separately and allowing the expression to occur. The host organism, which may be of eukaryotic or prokaryotic origin is grown under conditions ensuring expression of fused sequences. The fused polypeptide is then purified and a polypeptide of the invention separated from its fusion partner using a suitable method. The fusion polypeptide may in a particular 35

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embodiment of the invention still be capable of binding to MSH or an analogue thereof.

The second polypeptide to which a polypeptide of the invention is fused may in one particular embodiment of the invention be a DNA fragment encoding a diphtheria toxin, a staphylococcus protein, a ricin toxin, Pseudomonas endotoxin, abrin or fungal ribosome-inactivation proteins (RIP). In other embodiments of the invention the second DNA fragment may encode a subsequence of a melanotropic hormone receptor, an MSH receptor or an ACTH receptor.

The fusion polypeptides of the invention may be modified as well as other polypeptides of the invention, e.g. they may be glycosylated, coupled to a carbohydrate or lipid moiety, contain a palmitoyl anchor or a part thereof bound to a solid support and be provided with a detectable label.

The present invention also relates to a polypeptide of the invention in substantially pure form and to a method of producing the polypeptide. The method of producing a polypeptide of the invention comprises the following steps:

- 20 (a) inserting a DNA fragment of the invention in an expression vector,
  - (b) transforming a suitable host organism with the vector produced in step (a),
- (c) cultivating the host organism produced in step (b) under suitable conditions for expressing the polypeptide,
  - (d) harvesting the polypeptide, and
  - (e) optionally subjecting the polypeptide to posttranslational modifications.

40 A DNA fragment or an analogue or subsequence thereof of the invention encoding a polypeptide of invention can be modified before or after it has been inserted into the vector or organism for expression. The polypeptide product may also be subjected to modification. The modification may comprise substitution, addition, insertion, deletion or rearrangement of one or more nucleotides and amino acids in the DNA and polypeptide, respectively. The term "substitution" is intended to mean the replacement of one or more nucleotides 10 or amino acids in a DNA fragment or polypeptide of the invention. The term "addition" means addition of one or more nucleotides and amino acids at either end of a DNA fragment/polypeptide of the invention or a part of them. Insertion is intended to mean the introduction of one or more nucleotides and amino acids in a DNA fragment or polypeptide of the invention or a part of them. Deletion is intended to mean the removal of one or more nucleotides and amino acids from a DNA fragment or polypeptide of the invention or from a part of them. Rearrangement is intended to indicate that one 20 or more nucleotides or amino acids have been exchanged within the DNA or polypeptide sequence, respectively. The DNA fragment may, however, also be modified by mutagenesis either before or after inserting it in the organism. A DNA or pro-

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The polypeptide produced as described above may be subjected to posttranslational modifications such as thermal treatment, chemical treatment (formaldehyde, glutaraldehyde etc.) or enzyme treatment (peptidases, proteinases and protein modification enzymes). The polypeptide may be processed in a different way when produced in an organism as compared to its natural production environment. It may or may not be advantageous to remove or alter the processing characteristics caused by the host organism in question.

tein sequence of the invention may be modified in such a way

that it does not lose any of its biophysical, biochemical or

all) or all of such properties (one and/or all).

biological properties, or part of such properties (one and/or

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When a polypeptide according to the invention is produced in a prokaryotic organism such as a bacterium, a useful post-translational modification may be refolding of the peptide in order to obtain the peptide in a native and functional form due to the fact that peptides produced this way are often found as insoluble non-functional inclusion bodies inside the microorganism. The refolding of such peptides of such inclusion bodies are traditionally refolded by denaturing the peptide followed by a gradual continuous renaturation.

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The term "truncated" polypeptide refers to a polypeptide deleted of one or more amino acids eventually resulting in changing of the properties of the polypeptide, such as e.g. solubility. In a further meaning, the term "truncated" polypeptide refers to a mixture of polypeptides all derived from one polypeptide or expressed from the coding fragment(s) encoding said polypeptide.

Subsequent to the expression according to the invention of the polypeptide in an organism or a cell line, the polypeptide can either be used as such or it can first be 20 purified from the organism or cell line. If the polypeptide is expressed as a secreted product, it can be purified directly. If the polypeptide is expressed as an associated product, it may require the partial or complete disruption of the host before purification. Examples of the procedures employed for the purification of polypeptides are: (i) im-25 munoprecipitation or affinity chromatography with antibodies, (ii) affinity chromatography with a suitable ligand, (iii) other chromatography procedures such as gel filtration, ion exchange or high performance liquid chromatography or derivatives of any of the above, (iv) electrophoretic procedures like polyacrylamide gel electrophoresis, denaturating polyacrylamide gel electrophoresis, agarose gel electrophoresis and isoelectric focusing, (v) any other specific solubilization and/or purification techniques.

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Also, preparation of polypeptides of the invention may be performed by the well known methods of liquid or solid phase peptide synthesis utilizing the successive coupling of the individual amino acids of the polypeptide sequence. Alternatively, the polypeptide can be synthesized by the coupling of individual amino acids forming fragments of the polypeptide sequence which are later coupled so as to result in the desired polypeptide. These methods thus constitute another interesting aspect of the invention.

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10 Very important and thus constituting important aspects of the invention are various methods of regulating the activity exerted by an MSH receptor. This activity which has been described above in details may have important implications for the various disease conditions connected to the MSH receptor and for the various other biological functions. Thus, methods for preventing or stimulating the binding of the MSH receptor to various molecules constitute important aspects of the invention.

One of such aspects of the invention relates to a method of 20 preventing or stimulating the coupling of the described MSH receptor to its guanine nucleotide binding protein comprising using a method wherein a ligand is bound to an epitope of the receptor which normally interacts with the G-protein, in particular one or several of the extracellular loops and/or the transmembrane sequence and/or the cytoplasmic loops and/or the C-terminal sequence, thereby inhibiting or stimulating the coupling between the G-protein and the receptor. Thus, in one aspect this method comprises administering a substance to an animal, in particular a human, which substance in advance has been found to bind to a polypeptide 30 having the amino acid sequence shown in SEQ ID NO: 2 or 16 or a subsequence comprising any of the subsequences encoding one or several of the extracellular loops and/or the transmembrane sequence and/or the cytoplasmic loops and/or the C-terminal sequence or a combination thereof. 35

In this context a "ligand" will refer to a substance, which may be natural or synthetic, and which will bind in a preferably reversible, but also possibly irreversible manner to the MSH receptor.

5 Another method according to the invention of preventing or stimulating the binding of MSH or similar peptides or a Gprotein to the described MSH receptor comprises administering a substance to an animal, in particular a human, which substance in advance has been found to bind to a polypeptide 10 having the amino acid sequence shown in SEQ ID NO: 2 or 16 or an analogue or subsequence thereof comprising any of the subsequences encoding one or several of the extracellular loops and/or the transmembrane sequence and/or the cytoplasmic loops and/or the C-terminal sequence or a combination 15 thereof, so as to occupy the binding site of the receptor using an agonist, an antagonist, a blocker or a substance such as a derivative of MSH having a structure similar to MSH, and optionally thereby preventing or stimulating the generation of second messenger elements. The substance may be a synthetic ligand such as a peptide, an organic compound or 20 an antibody capable of binding to the receptor or a part thereof. The antibody may be a monoclonal or polyclonal antibody.

A method of reducing or increasing the binding affinity of
the MSH receptor is an interesting aspect of the invention
and may be obtained by the use of allosteric modulation.
Further a way of preventing the coupling of the MSH receptor
to its guanine nucleotide binding protein according to the
invention is to reduce the production of the MSH receptor.

This may be obtained by using antisense oligotherapy wherein
a DNA or RNA fragment complementary to at least part of the
mRNA corresponding to a polypeptide of the invention or an
analogue thereof may be effective in arresting the translation of the polypeptide in human cells and thereby inhibiting
the synthesis of MSH receptor polypeptide.

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A method for increasing the production of the MSH receptor may be receptor upregulation. Also a method of decreasing or increasing the generation of second messenger elements and/or increasing the production of the MSH receptor and/or optionally increasing or decreasing the binding affinity of MSH to the MSH receptor is part of the invention. The method comprises administering to an animal, in particular a human, a medicament which is or becomes bound to a substance, which substance in advance has been found to bind to a polypeptide having the amino acid sequence shown in SEQ ID NO: 2 or 16 or an analogue or subsequence comprising any of the subsequences encoding one or several of the extracellular loops and/or the transmembrane sequence and/or the cytoplasmic loops and/or the C-terminal sequence or a combination thereof.

15 A substance as the above indicated could be chosen by employing the methods described below for identifying substances
which can prevent or stimulate the effect exerted by MSH
receptors.

A method for internalization of an MSH receptor, thereby

making it unavailable for the binding of the hormone, constitutes another aspect of the invention. The method comprises using a substance which in advance has been found to bind to a polypeptide having the amino acid sequence shown in SEQ ID NO: 2 or 16 or a subsequence comprising any of the subsequences encoding one or several of the extracellular loops and/or the transmembrane sequence and/or the cytoplasmic loops and/or the C-terminal sequence or a combination thereof, such as a substance or a modified form of the MSH which is either able or unable to initiate the normal processes activated by the hormone but which substance causes internalization of the receptor. This method may also be used to regulate the effect exerted by the MSH receptor.

By using radioligand binding techniques and expressed MSH receptor protein and peptides the binding affinities of substances (ligands) for the MSH receptors may be measured. Such

45 measurements are typically performed in the screening of novel drugs (synthetic or natural) with potential activity on MSH receptors. In pharmacological terms such drugs may act as agonists, partial agonists or as antagonists at the MSH 5 receptor. All types of such substances are testable using the method according to the invention. The substances/ligands

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acting on the MSH receptor (or its derivative) can also be coupled to toxic agents (toxins, radionuclides) aiming to destroy the MSH receptor (or the MSH receptor derivative) bearing organisms or cells. 10

A very important aspect of the invention relates to methods for identifying substances which may be used for preventing or stimulating the effect exerted by an MSH receptor such as the generation of a second messenger element in a cell, such 15 as a mammalian cell, in particular a human cell. Inhibiting the binding of MSH to the MSH receptor is one way of achieving this, and therefore methods for identifying substances which are capable of binding to the MSH receptors are very important aspects of the invention. The methods involve various methods of assessing the capability of the substance in question to compete with the binding of MSH to the MSH receptor. The substance may prevent this binding by blocking the MSH binding site on the MSH receptor resulting in blocking of the effects exerted by the MSH receptor upon binding of MSH. In another embodiment, the substance may be a substance with optionally increased binding capacity to the MSH receptor (compared to MSH) and which is in addition capable of activating the effects exerted by the MSH receptor.

One embodiment of this aspect of the invention is to incubate an MSH receptor protein or its analogue, obtained as described in example 3, with radioactively labelled MSH or MSH analogue together with the test substance. Depending on the binding activity of the test substance the amount of labelled MSH or MSH analogue becoming bound to the MSH recep-35 tor will vary. Test substances with high binding affinity for the MSH receptor will exclude the binding of the labelled MSH WO 94/04674
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or MSH analogue at lower concentration than test substances with lower binding affinity. Separation of bound versus free labelled MSH or MSH analogue is accomplished using techniques such as filtration, centrifugation, superflow or chromatogra-5 phy. Measurement of radioactivity either retained on the receptor or being present in the solution separated from the receptors is made using standard nuclear counting. In another variant of this embodiment of the invention, the amount of MSH or MSH analogue being bound to the receptor or being 10 present in the solution separated from the receptors is detected by using any other suitable detection system capable of detecting MSH or MSH analogue. Examples of such detection systems are immune assays such as radio immune assay and ELISA (Enzyme linked immune sorbent assay), immune fluo-15 rescence assay, UV light absorption spectrometry or fluorescence emission spectrometry.

In another embodiment of this aspect of the invention, the amount of test substance bound to the MSH receptor is indirectly and/or approximately estimated by measuring the 20 alteration in the degree of interaction of the MSH receptor with a G-protein caused by the binding of test substance to the MSH receptor. In a variant of such a test system, the effect of the test substance is studied alone. In another variant of such an assay, the ability of the test substance 25 to compete for MSH or MSH analogue is studied by the simultaneous addition of test substance and MSH or MSH analogue. The degree of activation of the G-protein by the MSH receptor caused by the test substance, the MSH or the MSH analogue can be measured directly by e.g. measuring the GTPase activity of 30 the system using previously described methods (Aktories and Jakobs 1981; Vachon et al. 1986), or by using other suitable methods. Alternatively the degree of activation of the system may be studied indirectly by measuring other biochemical or physiological parameters which may become altered as a consequence of the primary interaction of the ligand(s) with the 35 MSH receptor. Examples of measurable entities in this context are adenylate cyclase activity, cAMP-levels, skin pigmentation, tyrosinase activity and [35]methionine incorporation

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(Burchill et al. 1990).

In yet another embodiment of the invention, which is particularly useful if the ligand is a macromolecule, such as when 5 the ligand is an antibody, the detection of the binding of the ligand to the receptor may be done using other approaches. In one variant of this embodiment, the ligand is incubated with a substantially pure preparation of the MSH receptor or its analogue, the latter which has been tagged 10 with a suitable molecule which, after separation of bound versus free MSH receptor, will allow the detection of the MSH receptor ligand complex by e.g. nuclear counting, colour, fluorescence or enzymatic activity. Separation of bound and free ligand may, for example, be accomplished by adding a second antibody which is directed towards the ligand thereby forming a precipitate of the ligand-receptor complex. In other variants of the invention, a substantially pure preparation of the MSH receptor is attached to a solid support. The ligand is then incubated with the solidified receptor 20 whereafter detection of the amount of ligand bound to the receptor may be done using conventional ELISA or using any similar suitable approach.

It will be understood that the ligands described herein may be provided with a detectable label. The ligands themselves can be macromolecules, such as monoclonal or polyclonal anti-25 bodies or they may be substances of natural or synthetic origin which are able to bind to MSH receptor.

Using the above and similar approaches substances can be identified which can block the binding of the MSH receptor (or its derivative) by the receptor ligands. In the present 30 context the term "blocking of the MSH receptor (or its derivative) " means that the MSH receptor (or its derivative) is occupied by the substance so that the receptor ligands cannot bind the MSH receptor (or its derivative) or that the MSH

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receptor ligands are capable of binding to the MSH receptor but unable to activate the MSH receptor.

It will be understood that methods similar to those mentioned above for identifying substances which bind to an MSH receptor can be used for identifying substances which bind to other melanotropic hormone receptors. Since the DNA fragments of the invention which have the nucleotide sequences shown in SEQ ID NO: 5 and 9, coding for the polypeptides of the invention shown in SEQ ID NO: 6 and 10 may be derived from other melanotropic hormone receptors, the binding properties of these melanotropic hormone receptors may be of great importance. Thus, important aspects of the invention are methods as the above-mentioned for identifying substances which are capable of binding to melanotropic hormone receptors.

It is of course also important to note, that the DNA fragments having the nucleotide sequence shown in SEQ. ID NO: 5, 7 and 9 can be modified in the same manner as other DNA fragments of the invention, and thus, all disclosure in the present specification relating to the modifications of the 20 DNA fragments having the nucleotide sequence shown in SEQ. ID NO: 1 and 15 applies analogously or mutatis mutandis to modifications of the DNA fragments having the nucleotide sequence shown in SEQ ID NO: 5, 7 and 9. Also, as aspects of the invention, the DNA fragments having the nucleotide sequences shown in SEQ ID NO: 5, 7 and 9 can be used in the 25 same manners as described in the present specification for the DNA fragments having the nucleotide sequences shown in SEQ ID NO: 1 and 15. Likewise, the polypeptides having the amino acid sequences SEQ ID NO: 6, 8 and 10 can be modified in the same manner as the polypeptides shown in SEQ ID NO: 1 30 and 15, and thus, all disclosure in the present specification relating to the modification of the polypeptides having the sequences shown in SEQ ID NO: 6, 8 and 10 applies analogously or mutatis mutandis to modifications of the polypeptides shown in SEQ ID NO: 1 and 15. Also, as aspects of the inven-35 tion, the polypeptides having the amino acid sequences shown

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in SEQ ID NO: 6, 8 and 10 can be used in the same manners as described in the present specification for the polypeptide having the amino acid sequence shown in SEQ ID NO: 1 and 15.

The polypeptide with the amino acid sequence SEQ ID No: 2 has been identified as an MSH receptor, based on the results of such binding experiments. A series of POMC (pro-opiomelano-cortin) derived peptides showed differential potencies in inhibiting  $^{125}\text{I-NDP-MSH}$  binding to MSH receptor coding fragment transfected COS-7 cells. The potency order was NDP-MSH  $(\text{K}_{\underline{i}} = 23 \pm 0.5 \text{ pM}) > \alpha\text{-MSH} \ (\text{K}_{\underline{i}} = 92 \pm 19 \text{ pM}) > \text{ACTH} \ (1\text{-}39)$   $(\text{K}_{\underline{i}} = 170 \pm 37 \text{ pM}) > \beta\text{-MSH} \ (\text{K}_{\underline{i}} = 449 \pm 74 \text{ pM}) > \gamma\text{-MSH} \ (\text{K}_{\underline{i}} = 1010 \pm 200 \text{ pM}). \ \text{ACTH} \ (4\text{-}10) \ \text{showed} \ \text{very low binding affinity}$   $(\text{K}_{\underline{i}} = 22,400 \pm 7200 \text{ pM}), \ \text{whereas the non-melanotropic POMC}$   $\text{peptide } \beta\text{-endorphin showed no affinity for the expressed MSH}$ 

Moreover, in addition the polypeptide with the amino acid sequence shown in SEQ ID NO: 16, referred to as\_the MC-2 receptor, has been identified as an MSH receptor and/or MSH receptor subtype based on the result of such binding experiments. A series of POMC derived peptides showed the following differential potencies in inhibiting 125I-NDP-MSH binding to MC-2 receptor coding fragment transfected COS-7 cells: NDP-MSH (Ki = 5.18  $\pm$  0.54 nM) >  $\alpha$ -MSH (Ki = 928  $\pm$  314 nM) = ACTH (1-39) (Ki = 929  $\pm$  389 nM) >  $\beta$ -MSH (Ki = 1.75  $\pm$  0.67  $\mu$ M) >  $\gamma$ -MSH (Ki = 3.45  $\pm$  0.88  $\mu$ M). The non-melanotropic POMC peptide  $\beta$ -endorphin showed no affinity for the expressed MC-2 receptor.

The above mentioned binding experiments can be done using whole animal systems, human clinical trials, a tissue specimen, a microorganism and/or a cell, in particular a cell line expressing the said receptor protein or its analogue. It can also be achieved using the purified protein of the invention. The purified protein can be used in a soluble form or in the solid phase being attached to a suitable matrix.

Drugs can be designed so as to act on very specific parts of a polypeptide of the invention. Drugs can be acting on either only the regions of or within the extracellular loops or transmembrane segments. In either case it may be affecting the binding of the natural ligands to the MSH receptor or its derivatives. Specific drugs can also be directed towards the regions of intracellular loops. Such drugs could be affecting the coupling of the MSH receptor or its derivatives to the

intracellular systems like the G-proteins. Such drugs could

10 also be affecting the G-proteins so that they cannot couple

to the MSH receptor or its derivatives.

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Various ways of treating the disease conditions wherein an MSH receptor is involved are provided by the present invention. These diseases include MSH receptor expressing disease condition such as melanoma, skin cancer, pyretic condition, inflammatory condition, nociceptive condition, catatonic condition, impaired memory condition, reduced or increased skin tanning and/or pigmentation conditions, epilepsy. The invention also includes a method to improve nerve repair, muscle reinnervation and/or neuron growth.

Due to its central nervous system localization the MC-2 receptor is in particular a target for drugs used in treating conditions such as pain, pyretic, catatonic and impaired memory conditions. Moreover, due to peripheral localization of the MC-2 receptor it is an interesting target for the anti-inflammatory drugs. The MC-2 receptor is also an interesting target for drugs improving growth and/or regeneration and/or repair of neurons being damaged due to disease and/or toxic influence and/or age and/or by other condition being associated with or leading to neuron damage. Moreover, due its ability to improve muscular reinnervation the MC-2 receptor is an interesting target for drugs treating condition of impaired muscle innervation. In addition due to its central nervous system localization the MC-2 receptor is also an important target for drugs used in the treatment of epilepsy.

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Thus, the invention relates to a method of targeting a cell that contains an MSH receptor on the surface with a medicament comprising administering the medicament in the form of a substance that binds to the MSH receptor. In a particular 5 embodiment of the invention, the medicament may be attached to a substance such as an antibody or a part thereof or be a molecule of natural or synthetic origin having affinity for the MSH receptor. The medicament may be a radionuclide or a toxin or any other molecule of natural or synthetic origin. 10 The use of an antibody such as a monoclonal antibody as a substance to which a medicament is bound comprises an important aspect of the invention. The antibody could be tagged with toxin or radioactivity for diagnostic or therapeutic purposes. Such an approach is expected to be superior to the 15 MSH toxin conjugates mentioned above, because of the expected high avidity and specificity of such MSH receptor antibodies. Moreover, the MSH toxin conjugates might induce untoward effects by virtue of their potential hormonal activity. The use of toxin or radiation coupled monoclonal antibodies 20 against the MSH receptor may prove to be a very attractive approach as the MSH receptor is the most common and most specific component of the melanoma cells.

In another embodiment of the invention the substance is a natural or synthetic organic compound, or a peptide or deri-25 vative thereof, that binds to the receptor or an epitope thereof and which optionally may become discovered by using methodology described in the present application. Such a substance may in particular be a synthetic and/or a natural compound which have or do not have any structural resemblance 30 to MSH. Such substances are typically composed of one or two or several aromatic and/or non-aromatic rings and/or heterocyclic rings, with side chains appropriately attached and may in addition have chains interconnecting the ring structures. Such substances vary considerably in their structure and there exist several different classes of such sub-35 stances which is due to the fact that they either bind to the same or the differing epitopes of the MSH receptor. Some of

these substances share partly or totally the same binding epitope on the MSH receptor as the MSH peptide, whereas other substances bind to other and/or partially other epitopes of the MSH receptor. Some of these substances have the ability to mimic the action of MSH in that when they bind to the receptor they cause the same effects in a cell and/or organ and/or tissue as when MSH binds to the MSH receptor. Other substances by contrast have the ability to prevent the action of MSH on the receptor by their binding to the MSH receptor.

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In a very important aspect of the invention, a lipid soluble form of an MSH receptor may be used in the treatment of an animal, in particular a human, by administering this form to the animal.

Also in a very important aspect of the invention, conditions caused by MSH receptor deficiency or impaired MSH receptor function in an animal, in particular a human may be treated by introducing a DNA fragment encoding an active form of an MSH receptor. One such condition which may be treated by the present invention is tyrosinase-positive albinism.

As stated previously, endogenous and exogenous melanotropins are suggested to enhance human cutaneous pigmentation in vivo (Levine 1991; Mulligan et al. 1982; Lerener et al. 1961). A treatment which produces tanning without sun exposure will be helpful to people who tan poorly and sunburn easily. Increased melanin in the skin might afford these people protection against ultraviolet light and thus put them at low risk for skin cancer.

The use of MSH and other melanotropins may be an effective and safe means of achieving skin darkening without harmful excess sun exposure. In addition, the resultant increased skin pigmentation might provide protection against the effects of subsequent sun exposure. In patients with tyrosinase-positive albinism, the molecular machinery to make melanin is present, but functions suboptimally (King et al.

1988). Perhaps, by acting on MSH receptor melanotropins could in these patients stimulate tyrosinase resulting in an increase in pigmentation. This may afford these individuals protection from ultraviolet light while improving their appearance and social acceptance.

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In the present context melanotropin is intended to be a substance that is binding to an epitope of an MSH receptor and thereby e.g. induces a similar skin tanning effect as MSH or any other desirable effect similar to that of MSH. Such a substance may be found using the methodology described in the present application and may in particular be a synthetic substance which have or do not have any structural resemblance to MSH.

Also, the invention relates to a method for increasing the
melanin content of the skin in an animal, in particular a
human, comprising using substances that are active through an
MSH receptor. Thus, the skin tanning may be obtained without
or with reduced exposure to sunlight which will make it
possible to avoid sunburns, which is most desirable as
already explained above.

Detection of the MSH receptors of the invention is important in various diagnostic aspects of the invention and may facilitate the diagnosis of various of the disease conditions associated with a content of MSH receptors in the tissue that is higher than normally found in said tissue and improve the prognosis of some of the diseases such as melanoma and skin cancer. Especially important aspects of the invention are the use of the detection of MSH receptor in MSH receptor expressing diseases, such as melanoma or skin cancer, in assessing the prognosis and/or guidance for further treatment.

Thus, the invention relates to a method of diagnosing an MSH receptor expressing disease condition such as melanoma or skin cancer comprising targeting a cell containing an MSH receptor on the surface with a diagnostic agent capable of

binding to the MSH receptor, which diagnostic agent can be detected following binding to the receptor. The diagnostic agent may be administered bound to a substance that binds to

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MSH receptor.

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5 The diagnostic agent may be a radioactive substance, or may be linked to a radioactive substance. In other embodiments, the diagnostic agent may be a coloured or colour generating substance or linked to a colour or colour generating agent.

One diagnostic method of the invention is detecting an MSH

receptor in a biological sample, wherein the sample is
treated with a substance that binds to the MSH receptor, and
detecting or visualizing the presence of the bound substance.

In a particular interesting embodiment of the invention, the
substance is an antibody or a part thereof. The antibody may

be an antibody that distinguishes between possible different
forms of the MSH receptor. The antibody may be labelled with
radionuclide, or biotinylated or may be unlabelled and later
detected by immunostaining. An important method in connection
with this part of the invention is detection and/or measurement of the bound antibody by a method of the ELISA type or
by a method of the radioimmunoassay type.

The terms "a sample" or "a biological sample" as used herein are defined as a cell, a subcellular fraction, a cell fraction, a tissue sample, a cell culture, or a cell suspension.

In connection with the above, the invention also relates to polyclonal and monoclonal antibodies which are reactive with a polypeptide or an analogue or subsequence thereof of the invention. A detailed description of the various aspect of the invention involving antibodies and which constitutes parts of the invention is given herein.

In the present context the term antibody is understood as the whole antibody molecule or any fragments thereof. An antibody can be fragmented during and/or after the production. It can

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also be made in the fragmented form to begin with and used as such or used after joining different fragments.

The animal used for the preparation of antibodies to a polypeptide of the invention is preferably selected from the group consisting of rabbit, monkey, sheep, goat, mouse, rat, pig, horse and guinea pigs. The cells producing the antibodies may be spleen cells or peripheral blood lymphocytes.

The antibody or fragments thereof may be of a monospecific (polyclonal) kind. The monospecific antibody may be prepared by injecting a suitable animal with a substantially pure preparation of a polypeptide of the invention. This can be followed by one or more booster injections at suitable intervals before the first bleeding. The animals are bled about 5-7 days after each immunization. Antibodies may optionally be isolated from the serum using standard antibody purification techniques (Sambrook et al. 1989).

Using the sequence of SEQ ID NO: 2 polyclonal antibodies have been prepared by chemically synthesizing two peptides which had the amino acid sequences identical to amino acids 4-19 and 25-35 of SEQ ID NO: 2, respectively.

These two peptides were (separately) coupled to thyroglobulin and separately injected into rabbits in Freund's adjuvant. After four booster injection both the conjugates were found to have induced formation of sera in the rabbits which were highly reactive against MSH receptor containing cells. The details of the manufacturing of the anti MSH rabbit sera is given in Example 10.

A monoclonal antibody or fragments thereof may be raised against an essential component of an MSH receptor, i.e. an epitope. The monoclonal antibody may be produced using conventional techniques (Köhler et al. 1975) by use of a hybridoma cell line, or by clones or subclones thereof or by cells carrying genetic information from the hybridoma cell

line producing said monoclonal antibody. The monoclonal antibody may be produced by fusing cells producing said monoclonal antibody with cells of a suitable cell line, and cloning the resulting hybridoma cells producing said monoclonal antibody. Alternatively, the monoclonal antibody may be produced

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body. Alternatively, the monoclonal antibody may be produced by immortalizing an unfused cell line producing said monoclonal antibody. The monoclonal antibodies are ultimately harvested from the cell growth medium. Hybridoma cells used to make monoclonal antibody may be grown in vitro or in the body cavity of an animal. The monoclonal antibody or fragments thereof may also be made using the recombinant DNA techniques (Huse et al. 1989).

Monoclonal antibodies may also be made by immunizing the suitable animals with a unpurified preparation of an MSH receptor protein. The resulting hybridoma clones secreting monoclonal antibodies can be screened for their ability to block the binding of MSH or its analogue to the MSH receptor e.g. using the approach described in example 3.

The idiotypic (antigen binding) structure of the antibody is antigenic and can thus give rise to specific antibodies directed against the idiotypic structure. The antibodies raised against the idiotype are called the anti-idiotypic antibodies. Such antibodies may mimic the structure of the original antigen and therefore may function as the original antigen. Such antibodies may be able to substitute the original antigen (MSH receptor protein, polypeptides or their analogues) for a part or all of the functions, usability and properties of the original polypeptide of the invention.

Preferably the monoclonal antibodies or fragments thereof will be used in most cases but polyclonal antibodies or fragments thereof may also be used. Typical uses of MSH receptor antibodies are as follows:

For purification of proteins: The antibodies can be used to purify an MSH receptor or its derivatives from the biological

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samples, using the affinity chromatography or the immunoprecipitation techniques.

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For diagnosis and therapy: The monoclonal antibodies against an MSH receptor or its derivatives can be used in the diagnosis and therapy of disease conditions in animals and humans. The diagnostic and therapeutic antibodies may be valuable for the disorders of skin, like skin cancer generally described as melanoma. The finding that MSH receptor is consistently found on melanoma tissues supports this notion.

The diagnostic agent may be an antibody with the specificity for a polypeptide of the invention. The antibody can be coupled to another protein or a solid support and/or can be used in the agglutination tests or the colour developing tests. Such antibodies can also be used to quantitate the MSH receptor or its derivatives in biological samples using the standard histochemistry or immunochemistry techniques.

For toxin therapy: The specific monoclonal antibodies can be coupled to different toxins like ricin or diphtheria toxin. Generally the A-chain of the plant toxin ricin or the A-chain 20 of the diphtheria toxin is conjugated to the monoclonal antibody in order to assemble hybrid proteins which have a targeted cytotoxicity. Moreover, the toxin used may alternatively be selected from Pseudomonas endotoxin, abrin or fungal ribosome-inactivation proteins (RIP). In the present context, a hybrid between the monoclonal antibody against a polypeptide of the invention and a toxin moiety can be used to bring about the killing of the MSH receptor bearing cells in an organism. Moreover, in the present context the toxin is intended to mean any toxin that is suitable for the purpose of killing and/or damaging the cell wherein the MSH receptor is located. The toxic effect of the toxin may be brought about when the toxin is still conjugated with the antibody. However, more likely the toxin will be processed once the antibody-toxin conjugate has become attached to the MSH receptor bearing cell. Such processing may involve e.g. 35 internalisation of the antibody-toxin complex, cleavage of

toxin from antibody and transportation of the toxin within the cell to its site of action. The processing is being done by the natural machinery of the MSH receptor bearing cell and careful engineering of the properties of the antibody-toxin complex will maximise its toxicity by affording the most favourable processing pathway for the complex.

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In order to improve the clinical efficiency of the antibodytoxin complex the monoclonal antibody can be designed so as to reduce its size by e.g. utilising F(ab')2 or Fab' frag-10 ments instead of the whole intact antibody. The reduction in the size of the molecule will increase the ability of the antibody-toxin complex to diffuse from the blood to the site of the tumour cell. Moreover, selecting a small sized toxin for the conjugation to the antibody, such as e.g. fungal RIP, 15 will afford the same effect. Moreover, elimination of the Fc fragment, as is achieved with the use of F(ab')2 or Fab' fragments for toxin conjugation, will eliminate the possibility that the antibody-toxin conjugate will bind to cells containing Fc receptors thus minimising non-specific binding 20 of the toxin complex to other cells than MSH receptor bearing cells. Thus, this measure will increase the selectivity of the antibody-toxin complex and increase its cytotoxic effect since a higher dose will be possible to administer. The problem of inducing a humoral immune response in the patent to whom the antibody-toxin conjugate is administered may be minimised by prior and/or concomitant administration of a drug which suppress the immune response. Such drugs may e.g. be selected from cyclophosphamide, prednisone, azathioprine and/or cyclosporin. Moreover, another approach for the same 30 purpose is to administer a monoclonal antibody directed towards CD4 antigen. Yet, another approach for the same purpose is to carefully engineer the antibody-toxin complex to minimise its immunogenicity. Such engineering is afforded by eliminating the most immunogenic epitopes of the complex while still retaining its ability to bind to the MSH receptor with high affinity and retaining its desired toxic effect. Moreover, the engineering will also have the purpose to in59 crease the stability of the complex after it has been admini-

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crease the stability of the complex after it has been administered to the patient. Increased stability is essential to afford a good therapeutic effect.

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Increased clinical efficiency of the antibody-toxin complex 5 may also be afforded by concomitant administration to the patient of another agent which will potentiate the toxicity of the complex. The other agent may e.g. be cyclophosphamide, daunorubicin and/or interferon. Moreover, increased toxicity may be afforded by simultaneously utilising antibody-toxin 10 complexes where two and/or several toxins with different mechanism for their toxicity has been included. As an alternative the simultaneous administration of another antibodytoxin complex which is directed for yet another melanoma protein, with the anti MSH receptor antibody-toxin complex of the present invention, will afford increased clinical efficiency. Yet another measure to afford increased clinical efficiency of the antibody-toxin complex will be afforded by the additional coupling to the MSH receptor antibody-toxin complex a suitable radionuclide which by virtue of RIT 20 induces a cytotoxic effect. The approach of RIT for therapy of melanoma is described further below.

For radiodiagnosis and radiotherapy: The monoclonal antibodies against an MSH receptor or its derivatives can be used in the diagnosis and therapy of disease conditions in animals and humans. The diagnostic and therapeutic antibodies may be valuable for the disorders like skin cancer generally described as melanoma. The finding that MSH receptor is consistently found on melanoma tissues supports this notion.

The specific monoclonal antibodies can also be coupled to different radionuclides like, <sup>123</sup>I, <sup>125</sup>I, <sup>131</sup>I, <sup>99m</sup>Tc, <sup>111</sup>In, <sup>186</sup>Re, <sup>188</sup>Re, <sup>211</sup>At, <sup>212</sup>Bi, <sup>90</sup>Y, <sup>67</sup>Cu and <sup>67</sup>Ga. Such radiolabelled antibodies can be used for diagnosis (Radioimmunoscintigraphy) or therapy (Radioimmunotherapy).

In the present context a radionuclide is intended to mean any radionuclide that is suitable for the purpose of detecting the cell and/or cell cluster that is hosting an MSH receptor. Coupling of radionuclides to anti MSH receptor antibodies may 5 be afforded by a number of techniques. For 123I, 125I and/or 131 I coupling is afforded by well established chloramine-T, iodogen, lactoperoxidase and/or hydroxyperoxidase methods. Moreover, radioiodination may as an alternative be afforded using Bolton-Hunter reagent. Radiolabelling with e.g. 111In and/or 90Y and/or 67Cu may be afforded by the coupling of a bifunctional chelating agent to the antibody and then adding the radionuclide to the complex. By virtue of the chelating ability of the complex the radionuclide will become attached to the complex. Suitable chelators may be selected from eg. isothiocyanatobenzyl EDTA (CITC), diethylenetriaminepentaacetic acid (DTPA) and be coupled via the mixed anhydride or the cyclic anhydride (Hnatowich 1990). However, since such complexes may provide somewhat unstable chelation and moreover during their manufacture intra and intermolecular cross linking of antibodies, other chelators such as e.g. GYK-DTPA 20 or SCN-Bz-DTPA may be used as an alternative (Hnatowich 1990). Radiolabelling of 99mTc to the antibody may be afforded by using direct labelling techniques such as by reducing disulphide bonds on the antibody thereby providing sites for stable attachment of 99mTc. One method for this end is to use tin which will afford reduction of disulphide bonds and adding [99mTc]pertechnetate which also will provide 99mTc by reduction with the tin (Hnatowich 1990). In the present context tin may be provided in the form of e.g. stannous tartrate or any other form suitable for the purpose. As an 30 alternative to tin another suitable reducing agent may be used such as e.g. dithiotreitol and/or 2-mercaptoethanol. Moreover, [99mTc]-glucartate and/or [99mTc]-phosphonate may substitute for [99mTc]pertechnetate as source for 99mTc. Yet another approach for 99mTc labelling of anti MSH antibodies is to use chelators as was described above. For the purpose of 99mTc chelation a promising concept is to couple metallothioneins to the anti MSH receptor antibody since the pro-

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teins afford strong chelation of 99mTc. Still another chelator that may be employed is the diamide demercaptide chelator (Fritzberg et al. 1986).

Radioimmunoscintigraphy procedure is based on the fact that 5 the labelled antibody will recognize the MSH receptor (or its derivative) on the cells, normal or diseased, and that the antibody will not bind to the cells devoid of the MSH receptor (or its derivative). The ultimate quality of the scintigraphic examination is dependent on the absolute quantity of the MSH receptor (or its derivative) in the specimen under examination and the background activity. It is possible to detect tumours using this technique when the tumour to non-tumour signal ratios are 1.5/1 or higher. Imaging is initially generally performed as planar scintigraphic examination. Anatomical landmarks are indicated with a point source, separately recorded, and afterwards added with computer assistance. A Single Photon Emission Computed Tomography (SPECT) can be performed, acquiring data by a 360 degree rotation of the gamma camera around the object under 20 examination. Transverse, coronal, sagittal or oblique sections are then reconstructed using mathematical calculations. SPECT appears to improve sensitivity and requires low tumour/non-tumour signal ratio. In the present context the above mentioned approach can be used in the diagnosis of skin cancers and other disease conditions where MSH receptor (or its derivative) is expressed and can be approached by the monoclonal antibody against a polypeptide of the invention.

Radioimmunotherapy (RIT) for killing diseased cells by a toxic agent bound to a specific monoclonal antibody is a 30 promising concept. In order to be efficient RIT has requirements besides the tumour/non-tumour signal ratio. The amount of radioactivity has to be sufficient to eradicate tumour without giving a high radiation dose to the surrounding normal tissue. Furthermore, distribution of the labelled monoclonal antibody in the tumour has to be homogeneous, allowing radiation of all tumour cells. The biological half

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life of the monoclonal antibody in the tumour has to be long enough to allow the radionuclide to exert maximal radiation effects. In the present context, the above mentioned approach can be used in the therapy of skin cancers and other disease conditions where MSH receptor (or its derivative) is expressed and can be approached by the monoclonal antibody

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As ligand binding blockers: The prevention of the binding of an MSH receptor (or its derivative) by the receptor ligands can be suitably performed by the antibodies with the specificity for a polypeptide of the invention. In the present context the term "blocking of an MSH receptor (or its derivative)" means that the MSH receptor (or its derivative) is occupied by the antibodies so that the receptor ligands cannot activate the MSH receptor (or its derivative).

against a polypeptide of the invention.

In accordance with the above, the invention also relates to an antibody capable of binding to a polypeptide of the invention provided with a detectable label, and to a polypeptide of the invention provided with a detectable label. The polypeptide or the antibody may in some embodiments be coupled to a solid support. The support may be selected from the group consisting of plates, strips, beads, particles, films and paper, and the solid support may be of latex, polystyrene, polyvinyl chloride, polyolefin, nylon, polyvinylidene difluoride, cellulose, silicone or silica.

Other methods for detection and/or quantitation of an MSH receptor comprise detection of the DNA or RNA and such methods are preferably based on the principles of hybridization which have been described in details above. Thus, in one such aspect the invention relates to a method for detection and/or quantitation of the mRNA of an MSH receptor comprising extracting RNA from a biological sample such as a subcellular fraction, a cell, a tissue sample, a cell culture or a cell suspension and measuring the hybridization of said RNA to a labelled DNA fragment of the invention or a labelled RNA

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fragment which can be constructed from a DNA fragment of the invention. Also, methods for measuring RNA such as northern blot or dot blot may be employed. The hybridization may be performed in situ or a labelled antisense mRNA probe may be 5 used. In another embodiment detection and/or quantitation of the MSH receptor mRNA may be obtained by extracting RNA from cells or tissues and converting it into cDNA for subsequent use in the polymerase chain reaction (PCR). The PCR primer(s) may be synthesized based on a DNA fragment of the invention such as the DNA fragments shown in SEQ ID NO: 1, SEQ ID NO: 3 or SEQ ID NO: 4 or any other DNA fragment of the invention. This method for detection and/or quantification may be used as a diagnostic method for diagnosing an MSH receptor expressing disease condition such as melanoma, skin cancer, pyretic 15 condition, inflammatory condition, nociceptive condition, catatonic condition, impaired memory condition, reduced or increased skin tanning condition and/or pigmentation condition.

In another specific embodiment of the invention this method
for the detection of MSH receptor RNA and/or DNA is used as
production control in the breeding of animals for obtaining a
desired fur and/or skin colour in the animal. Animals for
which such a control of fur and/or skin colour is desired may
be selected from e.g. mammals and reptiles and may in particular e.g. be a snake, alligator, crocodile, mink, fox,
hamster or chinchilla.

In yet a further embodiment of the invention the MSH receptor coding fragment(s) or a subsequence thereof is being analyzed in an animal by using e.g. cloning or PCR as described above. The thus obtained DNA and/or cDNA is subjected to sequence analysis using known methodology with the purpose of detecting a specific variant of an MSH receptor. The detection of such variants of MSH receptor may be desired e.g. in production control for the breeding of animals in order to obtain a desired skin and/or fur colour. Animals for which such a control of fur and/or skin colour is desired may be selected

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from e.g. mammals and reptiles and may in particular e.g. be a snake, alligator, crocodile, mink, fox, hamster or chinchilla.

In still a further embodiment of the invention a desired skin and/or fur colour of the animal is being obtained by introducing into the animal the desired variant of the MSH receptor by e.g. manufacturing a transgenic animal which will appropriately produce the MSH receptor variant. As an alternative the desired MSH receptor variant is being obtained by mutating a natural MSH receptor coding fragment in situ in the animal. Animals where the introduction of MSH receptor variants are desired may be selected from e.g. mammals and reptiles and may in particular e.g. be a snake, alligator, crocodile, mink, fox, hamster or chinchilla.

In the present context an MSH receptor variant is intended to mean a homologue and/or analogue of SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 15 or SEQ. ID NO: 16.

The lack of detailed structural information at the atomic level about the tertiary structure of the MSH receptor (a member of the G-protein coupled receptor family) continues to hamper efforts to understand receptor function at the molecular level. A polypeptide of the invention can be produced in large quantities using protein purification methods, and/or recombinant DNA techniques and/or synthetic chemistry approach. The polypeptide of the invention can then be cry-25 stallized. Crystallization, the ordered packing of molecules in three dimensions, relies on achieving the right balance of attractive and dispersive or repulsive forces between protein molecules in solution. To solubilize an integral membrane 30 protein, like an MSH receptor or its derivatives, it is necessary to replace the natural phospholipid environment by the detergent. The resultant shell of detergent around the protein is considered to be the most important factor impeding the formation of ordered crystals. Removal of the detergent leads to the precipitation of the protein and renders it useless for the crystallographic studies.

The main problem in crystallisation of an MSH receptor is to find the ideal detergents which forms stable or semi-stable 5 complexes with the membrane section of the MSH receptor, where there are 7 transmembrane sections. It is also a prerequisite that the overall structure of the receptor is not affected too much, something which can be validated using functional assays such as e.g. radioligand binding as has 10 been described in the present application. Several classes of detergents, which are useful for crystallisation, are presently known and may be applied. It is a good approach for each of these detergents to obtain individual phase diagram for the solubility of the MSH receptor and analyze these diagrams in detail in order to find optimal conditions for 15 crystallisation. Using such approach it will be possible to find and refine a buffered solution or a mixture of detergents and salts to force the protein unit to arrange intermolecular contacts for crystal growth. By salts are in 20 the present context intended mono and/or divalent ions which may support the interaction of individual MSH receptor molecules to arrange in a crystal lattice. Moreover, crystallisation may be afforded by alternatively or in addition adding MSH or an MSH analogue which will bind to the receptor and increase the possible surface(s) for intramolecular contact. 25 In addition the solubility of the MSH receptor may be analyzed using temperature gradients. The initial aggregation of nucleus, indicating crystallisation, can be analyzed in combination with e.g. laser light scattering. To obtain crystals the approaches such as e.g. sitting and hanging drop 30 as well as micro-batch applications may be utilised. Improved crystals may be obtained using microgravity conditions. As a final approach heavy atom cluster can also be applied.

Crystal aggregates may be analyzed by subjecting them to e.g. synchrotron radiation at a suitable wave length, such as e.g. an 1 Å wave length or a wave length more or less than 1 Å,

and collecting data for radiation diffraction. The application of anomalous scattering to solve the phase problem in crystallography can be applied. Moreover, cooling of crystals a suitable temperatures such as e.g. -10°C and 4°C or the flash freezing of MSH receptor crystals, which are very radiation sensitive, may be applied. The solution of 3D structure of the MSH receptor from X-ray diffraction pattern may be afforded using well known computational techniques.

Although no G-protein coupled receptor (family to which the MSH receptor belongs) has yet formed crystals, two other 10 integral membrane proteins, the photosynthetic reaction centre (Allen et al. 1987) and bacteriorhodopsin (Henderson et al. 1990), have been successfully studied. Similar techniques as singly or in combination can be applied to the cry-15 stallization and/or atomic structure determination of the polypeptides of the invention. Moreover, other techniques aiming at the elucidation of the 3D structure of proteins are being rapidly developed. One such technique, which is already far advanced, is two-dimensional NMR (Wright 1989), as well as modern multidimensional NMR-techniques. In order to uti-20 lize such a technique for elucidation of MSH receptor 3D structure it is required to have sufficient amounts of pure MSH protein and then obtain the appropriate two-dimensional or multidimensional NMR data which is used along with the known primary amino acid sequence of the receptor applying 25 appropriate computational methods. In addition computational methods are also being developed aiming to elucidate the 3D structure of proteins in the computer. These methods are generally and collectively referred to as molecular modelling. It is predicted that once the 3D structure of one 30 member of the G-protein coupled receptor family is solved, it will be possible to rapidly solve the 3D structure of the other members provided that their primary amino acid sequences are known, by using one, two or several of above mentioned methods. This is due to the predicted high similari-35 ties in the 3D structure of these receptors. Successful elucidation of the 3D structure of other classes of proteins,

such as the elucidation of the 3D structure of renin based on its homology to trypsin (Radung 1988), has already been successfully achieved, and an analogous approach may be used to obtain the 3D structure of an MSH receptor. In such an 5 approach the backbone of the MSH receptor protein is aligned with the backbone of another G-protein coupled receptor using the most homologous parts of the amino acid sequences (subsequences) of the two proteins. In general only the transmembrane segments of the receptors are considered in this alignment. After that the MSH receptor has been aligned with the 10 backbone of the other G-protein coupled receptors refinement of the structure of the MSH receptor is being made by careful positioning of the transmembrane segments of the MSH receptor, e.g. involving the rotation and tilting of transmembrane segments, as well as the positioning of amino acid side chains, until eventual Wan der Waals overlaps has been eliminated. Moreover, further refinement of the structure is being made by finding positions of the amino acid side chains which will form suitable bonds, such as e.g. hydrogen bonds, in between different amino acids of the same and/or the 20 adjacent transmembrane segments. Finally further refinement of the structure is being made by minimizing the energy of the system using well known computational techniques. The energy of the system is usually calculated by approximate methods e.g. by using the Amber force field but also the more 25 exact methods of quantum mechanical calculations may be applied. Such computations are readily being made using commercially available computer programmes such as e.g. Hyperchem, Sybyl etc.

It is understood that once the atomic structure of one of the G-protein coupled receptors is known, it will be relatively easy to do the same for other members, including an MSH receptor, of this very important receptor family.

Knowledge of the atomic structure on the one hand will help to understand the receptors function in minute details and on the other hand will facilitate the improvement of the speci10

fic drug developments through computational and/or other suitable methods. Among the methods that can be applied are 3D graphical analysis of epitopes, the docking of ligands to potential epitopes of the MSH receptor and de novo design of substances in the computer.

Thus, the present invention also relates to the use of a polypeptide sequence of the invention for the deduction of three dimensional structure of an MSH receptor or an analogue thereof having MSH binding capacity for use in the design of a substance capable of binding to the MSH receptor.

It will be understood that the above disclosed DNA fragments can be used for finding and isolating other similar DNA fragments, i.e. DNA fragments of the invention, by employing techniques like PCR and hybridization. Also, it is possible to use the polypeptides of the invention for designing DNA probes to be used for such finding and isolation. Thus, using this approach oligonucleotide primers are deduced from the polypeptide sequences of the present invention using the universal genetic code. Such primers can be used to perform PCR to find and isolate other similar DNA fragments as described in Examples 1 and 4. Thus, such uses of DNA fragment and polypeptides of the invention also constitute important aspects of the invention.

## 5. Figure legends

- Pigure 1. Agarose gel electrophoresis analysis. Lane 1 molecular weight markers. Lane 2 PCR reaction as described in Example 1. Three DNA products can be seen at 705 bp, 501 bp and 372 bp. The band at 372 bp position was later used to clone the full length coding fragment for human MSH receptor.
- Figure 2. Northern blot analysis of the tissue distribution of the GE4 mRNA. Ten  $\mu$ g of poly(A)<sup>+</sup> RNA from different tissues, namely brain (lane 2), thymus (lane 3), parathyroid gland (lane 4), Parotid gland (lane 5), salivary gland (lane

6), adrenal gland (lane 7), testis (lane 8), liver (lane 9), lung (lane 10), heart (lane 11), spleen (lane 12), skeletal muscle (lane 13), intestine (lane 14), colon (lane 15) and WM 266-4 human melanoma cells (lane 16), was separated by electrophoresis and blotted onto a membrane. It was then hybridized to <sup>32</sup>P-labelled GE4 probe. As can be seen a positive band was seen only in lane 16 corresponding to the WM 266-4 human melanoma cells. The GE4 DNA was later found to be a part of the MSH receptor coding fragment. See Example 1 for details.

10 Figure 3. Plasmid map of pB-11D.

Figure 4. Plasmid map of pE-11D.

Figure 5. 125 I-NDP-MSH binding to the WM266-4 human melanoma cells.

Figure 6. Relative potencies of melanotropins for inhibiting

15 125I-NDP-MSH binding to COS-7 cells (available from ATCC)

transfected with MSH receptor cDNA. Competition curves are shown for non-labelled NDP-MSH (■), α-MSH (●), ACTH (1-39)

(□), β-MSH (▲), γ-MSH (▼), ACTH (4-10) (o) and β-endorphin

(Δ). Experimental procedures are as described in Examples 2

20 and 3. Each point represents the mean of quadruplicate determinations. Non-specific binding was less than 8 percent of the total binding.

Figure 7. Plasmid map of pB-MC-2.

Figure 8. Plasmid map of pE-MC-2.

Figure 9. Relative potencies of melanotropins for inhibiting 125I-NDP-MSH binding to COS-7 cells (available from ATCC) transfected with pE-MC-2 plasmid DNA. Competition curves are shown for non-labelled NDP-MSH ( $\blacksquare$ ),  $\alpha$ -MSH ( $\bullet$ ), ACTH (1-39) ( $\square$ ),  $\beta$ -MSH ( $\blacktriangle$ ),  $\gamma$ -MSH ( $\blacktriangledown$ ), and  $\beta$ -endorphin (o). Experimental procedures are as described in Examples 7 and 8. Each point

represents the mean of quadruplicate determinations. Non-specific binding was less than 8 percent of the total binding.

Figure 10. Agarose gel electrophoresis analysis of the PCR products generated from the human brain and melanoma cells

5 mRNA. Lane 1-molecular weight markers, Lane 2-human brain mRNA without reverse transcription, Lane 3-human brain mRNA after reverse transcription, Lane 4-human melanoma cells mRNA without reverse transcription, Lane 5-human melanoma cells mRNA after reverse transcription. A specific product at the expected position of 380 bp is seen only in the human brain sample after reverse transcription.

## **EXAMPLES**

## EXAMPLE 1

Molecular cloning and nucleotide sequencing of the MSH receptor cDNA

## Design of polymerase chain reaction (PCR) primers

The primers were designed by careful examination of the sequence homologies in the membrane spanning segments 3 and 6 of the earlier cloned G-protein coupled receptors. The design of the primers was not inclined towards any one receptor or any one class of the receptors. The primers were degenerate (redundancy=8 for primer from segment 3 and redundancy = 32 for primer from segment 6) and also had an inert nucleotide called deoxyinosine at places which otherwise would have become highly degenerate. The primers were chemically synthesized on a custom order basis by Symbicom AB (Tvistevägen 48, Umeå, Sweden). Primers were synthesized with EcoRI (segment 3) and BamHI (segment 6) linkers at the 5'-ends to facilitate the cloning.

The primer sequences are shown in SEQ ID NO: 3 and SEQ ID NO: 4.

## Polymerase chain reaction (PCR) on human genomic DNA

One  $\mu$ g of human genomic DNA (Purchased from Clontech Laboratories Incorporated 4030 Fabian Way, Palo Alto CA 94303 USA) was subjected to PCR using the above described primers. The PCR was done in a final volume of 50  $\mu$ l. It contained one  $\mu$ g of human genomic DNA, 10 mM Tris-HCl pH 8.3, 50 mM KCl, 1.5 mM MgCl<sub>2</sub>, 0.001% gelatin, 200  $\mu$ M of each deoxynucleotide, 1  $\mu$ M of each primer and 1 unit of the enzyme Taq DNA polymerase (enzyme was obtained from Perkin Elmer Cetus, 761 Main Av., Noewalk, CT 06859, USA).

The PCR thermal profile used was 93°C for 60 seconds, 55°C for 40 seconds and 72°C for 60 seconds for a total of 25 cycles, using a thermal Cycler (Hybaid, 111-113 Waldegrave road, Teddington, Middlesex, TW11 8LL, UK).

Ten percent of the reaction was analyzed by agarose gel electrophoresis, using the standard methods (Sambrook et al. 1989) (see Fig. 1). Three products were identified at approximately 705 bp, 501 bp and 372 bp. The 705 and 501 bp products were later identified as DNA for two of the already cloned receptors, but the 372 bp product was identified as a novel G-protein coupled receptor, and was the only one processed further.

# 25 Sequencing of the 372 bp product

The 372 bp product was cloned into the pGEM7zf(+) vector (Promega Corp. Madison, WI, USA) using the standard techniques (Sambrook et al. 1989) and one of the resulting plasmid termed as GE4 was sequenced using the chain termination method (Sanger et al. 1977). It was found to contain the sequence shown in SEQ ID NO: 11.

## Tissue distribution of the GE4 mRNA

Poly(A) + RNA was prepared using the oligo-dT purification scheme as described in standard protocols (Sambrook et al. 1989) from the following tissues: brain, thymus, parathyroid gland, parotid gland, salivary gland, adrenal gland, testis, liver, lung, heart, spleen, skeletal muscle, intestine, colon and WM 266-4 human melanoma cells (ATCC # CRL 1676). Ten  $\mu$ g of Poly(A) + RNA was subjected to electrophoresis through a 0.8% agarose-formaldehyde gel as described (Sambrook et al. 1989). The RNA was then blotted on to a Genescreen membrane 10 (New England Nuclear, USA), and cross linked to the membrane with UV light. The membrane was then placed in a sealed plastic bag containing 10 ml of prehybridization solution (50% formamide, 5X SSC, 5X Denhardt's solution, 0.1% SDS, 10mM Sodium phosphate pH 7.0, 10mM EDTA and 100  $\mu$ g/ml de-15 natured calf thymus DNA) at 42°C for 4 hours. The prehybridization solution was then replaced with the 10 ml of hybridization solution (10 ml prehybridization solution + 32P labelled GE4 DNA probe). The GE4 DNA was labelled with 32P using a commercial multiprime kit (Amersham, Aylesbury, Buckinghamshire, HP20 1BR, U.K.). The membrane was left in the hybridization solution for 12 hours at 42°C. The membrane was then washed in a solution of 0.1% SSC and 0.1% SDS at 60°C for 30 minutes, air dried and then exposed to autoradiographic film for 16 hours. See Fig. 2. A positive signal can only be seen in the lane #16, which is for human melanoma WM 266-4 cells.

# Construction and screening of the cDNA library from WM 266-4 cells

The WM 266-4 cells were obtained from ATCC, Bethesda, MD, USA. The cells were grown in the medium as advised by the ATCC. Poly(A)<sup>+</sup> RNA from these cells was made using the fast track mRNA isolation kit (InVitrogen corp., 3985 Sorrento Valley Blvd. #B, San Diego, CA 92121, USA). Five μg of this RNA was then used to make a random primed cDNA library in the

lambda gt11 vector using the materials and the conditions described by the manufacturer of the kit used (Amersham, Aylesbury, Buckinghamshire, HP20 1BR, U.K.).

Approximately 7x10<sup>5</sup> plague forming units from the unamplified 5 library were plated on the agar-LB plates (Sambrook et al. 1989), grown for 8 hours and were then transferred to Hybond-C filter discs (Amersham, Aylesbury, Buckinghamshire, HP20 1BR, U.K.). The DNA on the filter discs was then denatured and fixed as described (Sambrook 1989). The filter 10 discs were then placed in sealed bags (4 filters/bag) containing the prehybridization solution (6X SSC, 5X Denhardt's solution, 10 mM sodium phosphate pH 7.0, 1 mM EDTA, 0.5% SDS and 0.1 mg/ml of denatured Salmon testis DNA) for 6 hours at 60°C. The filters were then placed in the hybridization solution (prehybridization solution + 32P-labelled GE4 DNA probe), for 12 hours at 60°C. The GE4 DNA was labelled with 32p using a commercial multiprime kit (Amersham, Aylesbury, Buckinghamshire, HP20 1BR, U.K.). The filters were then washed at 65°C in a solution of 0.1% SSC and 0.1% SDS for 20 minutes, air dried and then exposed to the autoradiographic film for 24 hours. The positive plaques were picked and after repeating the screening for two more times a positive plaque designated 11D was isolated.

# Subcloning and sequencing of the 11D cDNA

25 A large scale lambda DNA preparation was made for the 11D clone as described (Sambrook et al. 1989). The insert was excised out with the EcoRI and HindIII enzymes. This took out all of the coding and 5'-untranslated sequences with part of the 3'-untranslated sequence. The EcoRI-HindIII fragment was cloned between the EcoRI and Hind III sites of the pGEM7Zf(+) vector (Promega Corp., Madison, Wisconsin, USA) using the standard methods described (Sambrook 1989). The resulting plasmid DNA (See Fig.3; pB-11D) was then transfected into competent DH5alfa E.Coli (BRL, 8400 Helgerman court, Gaithersburg, MD 20877, USA). Bacterial colonies were grown on

agar plates containing ampicillin. Individual colonies were picked in ampicillin containing 5 ml LB medium (Sambrook 1989) and grown overnight. Plasmid DNA was prepared as described (Sambrook 1989). Prepared plasmid DNAs were checked 5 for the presence of 11D cDNA in correct position. The plasmid constructs with 11D cDNA in correct position were then amplified further in 500 ml cultures. Plasmid DNA from such large scale preparations were prepared with Qiagen Kits (Qiagen Inc., 11712 Moorpark Street, Studio City, CA 91604, USA). Both the strands of DNA in the entire coding sequence and the 5'-untranslated region and a small portion of the 3'-untranslated region were sequenced by making the overlapping fragments. The method of sequencing was the chain termination method (Sanger et al. 1977). The cloned 11D cDNA 15 was found to have the nucleotide sequence shown in SEQ ID NO: 1 and was shown to contain 7 hydrophobic segments (corresponding to nucleotides 286-351, 394-465, 517-588, 640-711, 733-804, 898-972 and 997-1068 in SEQ ID NO: 1, respectively).

#### EXAMPLE 2

20 Cloning of the 11D cDNA into an expression vector

The expression vector pcDNAI (Invitrogen Corp., 3985 Sorrento Valley Blvd. #B, San Diego, CA 92121, USA), is a 4.2 kb multifunctional eukaryotic expression vector. It has the human CMV promoter and enhancer for high level expression.

The 11D cDNA was excised from pGEM7Zf(+) vector (described above in 5.1.6) with EcoRI and NsiI enzymes, and ligated into the same sites of pcDNAI vector. The resulting plasmid DNA (See Fig.4; pE-11D) was then transfected into competent MC1061/P3 E.Coli. Bacterial colonies were grown on agar plates containing ampicillin. Individual colonies were picked in 5 ml LB-ampicillin medium (Sambrook 1989) and grown overnight. Plasmid DNA was prepared as described (Sambrook 1989). Prepared plasmid DNAs were checked for the presence of 11D cDNA in correct position. The plasmid constructs with 11D

cDNA in correct position were then amplified further in 500 ml cultures. Plasmid DNA from such large scale preparations were prepared with Qiagen Kits (Qiagen Inc., 11712 Moorpark street, Studio city, CA 91604, USA).

## 5 EXAMPLE 3

Expression of the 11D cDNA and establishment of its identity

The WM266-4 human melanoma cells (from which the MSH receptor cDNA has been cloned) were grown under conditions described by ATCC. These cells were subjected to radioligand binding as described below for the transfected COS-7 cells. The WM-266-4 cells were shown to bind the <sup>125</sup>I-labelled NDP-MSH in a specific manner. (Fig. 5)

COS-7 cells were grown in Dulbecco's modified Eagle medium with 8% foetal calf serum and non-essential amino acids (Gibco/BRL, 8400 Helgerman Court, Gaithersburg,-MD 20877, 15 USA). Eighty percent confluent cultures were transfected with 1  $\mu$ g of pE-11D plasmid DNA and 40  $\mu$ g lipofectin (BRL, 8400 Helgerman Court, Gaithersburg, MD 20877, USA) in serum free medium. Five hours after transfection, serum containing medium was replaced, and cells were cultivated for 20 hours. 20 Cells were then scraped off, centrifuged, resuspended in serum containing medium, plated on 48 well plates, and allowed to grow for 24 hours. The cells were then washed with 0.3 ml of binding buffer (minimum essential medium with Earle's salts, 25 mM HEPES pH 7.0, 0.2% bovine serum albumin, 25 1 mM 1,10-phenanthroline, 0.5 mg per litre leupeptine and 200 mg per litre bacitracin) and then incubated at 37°C for 2 hours with 0.3 ml binding buffer containing 24,000 cpm of 125I-NDP-MSH and appropriate concentration of unlabelled peptides. NDP-MSH was labelled with 125 Iodine (see below for 30 details) to the specific activity of 8.6x104 cpm per Mol. The plates were then put on ice, cells washed with 0.3 ml of ice cold binding buffer and detached from plates with 0.3 ml of 0.1 N NaOH. Radioactivity was counted and data analyzed by

iterative, non-linear curve fitting programme suitable for radioligand binding analysis (see Fig. 6). A series of POMC (pro-opiomelanocortin) derived peptides (purchased from Saxon Biochemicals GmbH, Hannover, Germany) showed differential 5 potencies in inhibiting 125I-NDP-MSH binding to pE-11D transfected COS-7 cells. The potencies and reciprocals of binding affinities (Kis) were determined by testing several (e.g. 11 - 12) concentrations of every tested peptide and fitting the data for the counts found to be bound to the cells to the 10 four parameter logistic function using non-linear regression analysis using previously described methods (Bergström and Wikberg 1986). The Ki-values were then calculated from the IC-50 values estimated thus estimated by using the Cheng and Prusoff equation, as previously described (Cheng and Prusoff 15 1973). The potency order and  $K_i$  values found from the analysis were NDP-MSH ( $K_i = 23 \pm 0.5 \text{ pM}$ ) >  $\alpha$ -MSH ( $K_i = 92 \pm 19 \text{ pM}$ ) > ACTH (1-39) (K<sub>i</sub> = 170 ± 37 pM) >  $\beta$ -MSH (K<sub>i</sub> = 449 ± 74 pM) > $\gamma$ -MSH (K<sub>i</sub> = 1010 ± 200 pM). ACTH (4-10) showed very low binding affinity ( $K_i = 22,400 \pm 7200 \text{ pM}$ ), whereas the non-melanotropic POMC peptide  $\beta$ -endorphin showed no affinity for the expressed MSH receptor. These results conclusively prove that the cloned DNA of the invention is the MSH receptor cDNA.

## Iodination of NDP-MSH

Four μg of the peptide NDP-MSH was iodinated with 1 mCi of
125-iodine using the Iodobeads (Pierce, Rockford, IL, USA) in
100 mM sodium phosphate buffer (pH 6.5) for 10 minutes. The
Iodobead was then removed from the solution which was applied
to the C-18 reverse phase chromatography cartridge pre-equilibrated with 15% acetonitrile/0.05 M ammonium acetate pH

5.8. The cartridge was washed with 5 ml of the pre-equilibration buffer and then eluted at a flow rate of 1 ml/minute
using a peristaltic pump. The elution gradient was 15% to 35%
of acetonitrile containing 0.05 M ammonium acetate pH 5.8.
Fractions of 1 ml were collected and the radioactivity determined by counting 2.5 μl from each fraction on to a gamma
counter. Fractions 25 to 29 were pooled, dried under vacuum

and redissolved in 1 ml water. The radioactivity was counted and the specific activity was calculated.

## EXAMPLE 4

Identification of DNA sequences related to the cloned MSH receptor cDNA

Two PCR primers were designed based on the sequence of the cloned MSH receptor cDNA. Their nucleotide sequences are shown in SEQ ID NO: 13 and in SEQ ID NO: 14, respectively.

These primers were used to perform PCR on human genomic DNA
in exactly the same way as described in Example 2 except for
the thermal profile, which was 94°C for 30 sec, 45°C for 20
sec, 72°C for 20 sec for 5 cycles and then 94°C for 30 sec,
60°C for 20 sec, 72°C for 20 sec for 25 cycles. Ten percent
of the reaction was analyzed by agarose gel electrophoresis,
using the standard methods (Sambrook et al. 1989). The products obtained were cloned into the pGEM7zf(+) vector and
sequenced to completion. They were shown to have the
nucleotide sequences shown in SEQ ID NO: 5, SEQ ID NO: 7 and
SEQ ID NO: 9, respectively.

#### 20 EXAMPLE 5

Cloning of the 11D cDNA into a stable expression vector

## Cloning of the 11D cDNA

The expression vector pRC/CMV (Invitrogen corp., 3985 Sorrento Valley Blvd. #B, San Diego, CA 92121, USA), is a 4.5 kb multifunctional eukaryotic expression vector. It has the CMV promoter and enhancer for high level expression, and neomycin gene for selection of stable transfectants.

The 11D cDNA was excised from pcDNA I vector as described in example 2 with Hind III enzyme and ligated into the same site

of pRC/CMV vector. The resulting plasmid DNA was then transfected into competent INVαF' E. coli. Bacterial colonies were grown on agar containing ampicillin. Individual colonies were picked in 5 ml LB-ampicillin medium (Sambrook 1989) and grown overnight. Plasmid DNA was prepared as described (Sambrook 1989). Prepared plasmid DNAs were checked for the presence of 11D cDNA in correct position. The plasmid constructs with 11 D cDNA in correct were then amplified further in 500 ml cultures. Plasmid DNA from such large scale preparations were prepared with Qiagen Kits (Qiagen Inc., 11712 Moorpark Street, Studio City, CA 91604, USA).

# Stable expression of the 11D cDNA

COS-7 cells (available from ATCC) were grown in Dulbecco's modified Eagle medium with 8% fetal calf serum and non-essential amino acids (Gibco/BRL, 8400 Helgerman Court, Gaithersburg, MD 20877, USA) in serum free medium, five hours after transfection, serum containing medium was replaced, and cells were cultivated for 48 hours. At this time selection for the calls stably harbouring the pRC/CMV-11D plasmid was begun by growing the cells in neomycin (0.5 mg/ml) containing medium (selection medium). Selection procedure was continued for two weeks, replacing the selection medium every 4th day. surviving cells were collected and maintained in the selection medium. These cells constitute the stable cell line.

To investigate the binding properties cells were scraped off from the culture flask, centrifuged, resuspended in selection medium, plated on 48 well, and allowed to grow for 24 hours. The cells were then washed with 0.3 ml of binding buffer (minimum essential medium with Earle's salts, 25 mM HEPES pH 7.0, 0.2% bovine serum albumin, 1 mM 1,10-phenanthroline, 0.5 mg per litre leupeptine and 200 mg per litre bacitracin) and then incubated at 37°C for 2 hours with 0.3 ml binding buffer containing 24,000 cpm of <sup>125</sup>I-NDP-MSH and appropriate concentration of unlabelled peptides. NDP-MSH was labelled

with 125 Iodine as explained in example 3 to the specific activity of 8.6x104 cpm per Mol. The plates were then put on ice, cells washed with 0.3 ml of ice cold binding buffer and detached form plated with 0.3 ml of 0.1 N NaOH. Radioactivity was counted and data analyzed by an iterative, non-linear curve fitting programme suitable for radioligand binding analysis. A series of POMC (pro-opiomelanocortin) derived peptides (purchased from Saxon Biochemicals GMBH, Hannover, Germany) showed differential potencies in inhibiting 123I-10 NDP-MSH binding to the stable cell line. The potency order found from the analysis were NDP-MSH >  $\alpha$ -MSH > ACTH(1-39) >  $\beta$ -MSH >  $\gamma$ -MSH. ACTH(4-10) showed very low binding affinity, whereas the non-melanotropic POMC peptide  $\beta$ -endorphin showed no affinity for the expressed MSH receptor. These results conclusively prove that the cloned MSH receptor cDNA is produced and expressed in a stable cell line.

## EXAMPLE 6

Molecular cloning and nucleotide sequencing of the full length clone of G8 DNA

# 20 Screening of a human placental genomic library

A human genomic DNA library was purchased from Stratagene, USA. Approximately 7x10<sup>5</sup> plaque forming units from this library were plated on the agar-LB plates (Sambrook et al. 1989), grown for 8 hours and were then transferred to
25 Hybond-N filter discs (Amersham, Aylesbury, Buckinghamshire, HP20 1BR, U.K.). The DNA on the filter discs was then denatured and fixed as described (Sambrook et al. 1989). The filter discs were then placed in sealed bags (4 filters/bag) containing the prehybridization solution (6X SSC, 5X Denhardt's solution, 10 mM sodium phosphate pH 7.0, 1 mM EDTA, 0.5% SDS and 0.1 mg/ml of denatured Salmon testis DNA) for 6 hours at 60°C. The filters were then placed in the hybridization solution (prehybridization solution + <sup>32</sup>P-labelled G-8 DNA probe, as in SEQ ID NO: 7), for 12 hours at 60°C. The G-8

DNA was labelled with <sup>32</sup>P using a commercial multiprime kit (Amersham, Aylesbury, Buckinghamshire, HP20 1BR, U.K.). The filters were then washed at 65°C in a solution of 0.1% SSC and 0.1% SDS for 20 minutes, air dried and then exposed to the autoradiographic film for 24 hours. The positive plaques were picked and after repeating the screening for two more times a positive plaque containing a gene designated MC-2 was isolated.

# Subcloning and sequencing of the full length MC-2 DNA

A large scale lambda DNA preparation was made for the MC-2 clone as described (Sambrook et al. 1989). The insert was excised out with the Sac I enzyme. This took out a 2.4 kb fragment containing all of the coding and a parts of 5'- and 3'-untranslated sequence. The Sac I fragment was cloned in the Sac I site of the pGEM5Zf(+) vector (Promega Corp., Madison, Wisconsin, USA) using the standard methods described (Sambrook 1989). The resulting plasmid DNA pB-MC-2 (Fig. 7) was then transfected into competent DH5alfa E.Coli (BRL, 8400 Helgerman court, Gaithersburg, MD 20877, USA). Bacterial 20 colonies were grown on agar plates containing ampicillin. Individual colonies were picked in ampicillin containing 5 ml LB medium (Sambrook et al. 1989) and grown overnight. Plasmid DNA was prepared as described (Sambrook et al. 1989). Prepared plasmid DNAs were checked for the presence of MC-2 DNA in correct position. The plasmid constructs with MC-2 DNA in 25 correct position were then amplified further in 500 ml cultures. Plasmid DNA from such large scale preparations were prepared with Qiagen Kits (Qiagen Inc., 11712 Moorpark Street, Studio City, CA 91604, USA). Both the strands of DNA in the entire coding sequence and the 5'-untranslated region 30 and a small portion of the 3'-untranslated region were sequenced by making the overlapping fragments and primer walking. The method of sequencing was the chain termination method (Sanger et al. 1977). The cloned MC-2 DNA was found to contain the sequence shown in SEQ ID NO: 15.

#### EXAMPLE 7

Cloning of the MC-2 DNA into an expression vector

The expression vector pRC/CMV (Invitrogen Corp., 3985 Sorrento Valley Blvd. #B, San Diego, CA 92121, USA), is a 4.5 kb multifunctional eukaryotic expression vector. It has the human CMV promoter and enhancer for high level expression.

The MC-2 DNA (from nucleotide no. 520 to nucleotide no. 1620 from the seq. I.D. no. 10) was ligated between the HindIII and the XbaI sites of the pRC/CMV vector. The resulting plasmid DNA pE-MC-2 (Fig. 8) was then transfected into competent INVαF' E.Coli. Bacterial colonies were grown on agar plates containing ampicillin. Individual colonies were picked in 5 ml LB-ampicillin medium (Sambrook et al. 1989) and grown overnight. Plasmid DNA was prepared as described (Sambrook et al. 1989). Prepared plasmid DNAs were checked for the presence of MC-2 DNA in correct position. The plasmid constructs with MC-2 DNA in correct position were then amplified further in 500 ml cultures. Plasmid DNA from such large scale preparations were prepared with Qiagen Kits (Qiagen Inc., 11712 Moorpark street, Studio city, CA 91604, USA).

#### EXAMPLE 8

Expression of the MC-2 DNA and establishment of its identity

COS-7 cells were grown in Dulbecco's modified Eagle medium with 8% foetal calf serum and non-essential amino acids
(Gibco/BRL, 8400 Helgerman Court, Gaithersburg, MD 20877, USA). Eighty percent confluent cultures were transfected with 1 μg of pE-MC-2 plasmid DNA and 40 μg lipofectin (BRL, 8400 Helgerman Court, Gaithersburg, MD 20877, USA) in serum free medium. Five hours after transfection, serum containing medium was replaced, and cells were cultivated for 20 hours. Cells were then scraped off, centrifuged, resuspended in serum containing medium, plated on 48 well plates, and

allowed to grow for 24 hours. The cells were then washed with 0.3 ml of binding buffer (minimum essential medium with Earle's salts, 25 mM HEPES pH 7.0, 0.2% bovine serum albumin, 1 mM 1,10-phenanthroline, 0.5 mg per litre leupeptine and 200 mg per litre bacitracin) and then incubated at 37°C for 2 hours with 0.3 ml of binding buffer containing 24,000 cpm of 125I-NDP-MSH and appropriate concentration of unlabelled peptides. NDP-MSH was labelled with 125 Iodine (see below for details) to the specific activity of 8.6x104 cpm per Mol. The plates were then put on ice, cells washed with 0.3 ml of ice cold binding buffer and detached from plates with 0.3 ml of 0.1 N NaOH. Radioactivity was counted and data analyzed by iterative, non-linear curve fitting programme suitable for radioligand binding analysis (see Fig. 9). A series of POMC (pro-opiomelanocortin) derived peptides (purchased from Saxon 15 Biochemicals GmbH, Hannover, Germany) showed differential potencies in inhibiting 125I-NDP-MSH binding to pE-MC-2 transfected COS-7 cells. The potencies and reciprocals of binding affinities  $(K_is)$  were determined by testing several (e.g. 10 - 12) concentrations of every tested peptide and 20 fitting the data for the counts found to be bound to the cells to the four parameter logistic function using non-linear regression analysis using previously described methods (Bergström and Wikberg 1986). The Ki-values were then calculated from the IC-50 values estimated thus estimated by using 25 the Cheng and Prusoff equation, as previously described (Cheng and Prusoff 1973). The potency order and  $K_i$  values found from the analysis were NDP-MSH ( $K_i = 5.18 \pm 0.54 \text{ nM}$ ) >  $\alpha$ -MSH ( $K_i$  = 928 ± 314 nM) = ACTH (1-39) ( $K_i$  = 929 ± 389 nM) >  $\beta$ -MSH (K<sub>i</sub> = 1.75 ± 0.67  $\mu$ M) >  $\gamma$ -MSH (K<sub>i</sub> = 3.45 ± 0.88  $\mu$ M). 30 The non-melanotropic POMC peptide  $\beta$ -endorphin showed no affinity for the expressed MC-2 receptor. These results conclusively prove that the cloned MC-2 DNA of the invention is a new member of the melanotropic receptor family.

Iodination of NDP-MSH: Four mg of the peptide NDP-MSH was iodinated with 1  $\mu$ Ci of <sup>125</sup>Iodine using the Iodobeads (Pierce, Rockford, IL, USA) in 100 mM sodium phosphate buffer

(pH 6.5) for 10 minutes. The Iodobead was then removed from the solution which was applied to the C-18 reverse phase chromatography cartridge preequilibrated with 15% acetonitrile/0.05 M ammonium acetate pH 5.8. The cartridge was washed with 5 ml of the pre-equilibration buffer and then eluted at a flow rate of 1 ml/minute using a peristaltic pump. The elution gradient was 15% to 35% of acetonitrile containing 0.05 M ammonium acetate pH 5.8. Fractions of 1 ml were collected and the radioactivity determined by counting 2.5  $\mu$ l from each fraction on to a gamma counter. Fractions 25 to 29 were pooled, dried under vacuum and redissolved in 1 ml of water. The radioactivity was counted and the specific activity was calculated.

#### EXAMPLE 9

# 15 MC-2 RNA detection by PCR analysis

RNA from human brain tissue (purchased from Clontech, USA) and WM266-4 melanoma cells (made by Fast Track kit from Invitrogen Corp. USA) were reverse transcribed with Super-Script RNase H<sup>-</sup> reverse transcriptase (BRL, USA). PCR was performed on samples before and after reverse transcription to rule out the possibility of genomic DNA contamination in RNA preparations. Five µg of RNA was used for reverse transcription and then all of it was used as template in the first PCR. The first PCR was performed with primers (described below as number 1 & 2) specific for the 5'- and 3'-untranslated regions of the MC-2 DNA. Ten percent of the first PCR reaction was then subjected to a second PCR with primers (described below as number 3 & 4) specific for the coding region of the clone MC-2.

Primer 1: 5'-GGAAGCTTTCTTTGGTAGGCTG (SEQ ID NO: 17)

Primer 2: 5'-GGTCTAGAGCCACAGAGAGGAG (SEQ ID NO: 18)

Primer 3: 5'-CTGCATTTCTTGGATCT (SEQ ID NO: 19)

Primer 4: 5'-AAGCTGCACATGGATGC (SEQ ID NO: 20)

Both the PCRs were performed with Gene amplification kit
(Perkin Elmer Cetus, USA). The PCR thermal profile used was
93°C for 60 seconds, 55°C for 40 seconds and 72°C for 60
seconds for a total of 40 cycles. Fifty percent of the reaction was analyzed by agarose gel electrophoresis. The product
was seen at the expected 380 bp position (fig. 10).

## EXAMPLE 10

Development of polyclonal antibodies against the MSH receptor with polypeptide sequence according to SEQ ID NO: 2

The following two peptides, which were based on the polypeptide in SEQ ID NO: 2, were synthesized:

Peptide M1-Y, amino acids 4-19 of SEQ ID NO: 2:

Gln Gly Ser Gln Arg Arg Leu Leu Gly Ser Leu Asn Ser Thr Pro

15 Cys

Peptide M2-Y, amino acids 25-35 of SEQ ID NO: 2:

Gly Leu Ala Ala Asn Gln Thr Gly Ala Arg Cys

The peptides were synthesized commercially by the multiple peptide system, U.S.A. The synthetic peptides were conjugated to thyroglobulin (THY) by use of the MBS method. This method allows coupling of the free sulfhydryl group of the cysteine-containing peptide onto the carrier protein via the bifunctional crosslinker MBS.

## Immunization of rabbits

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Each of the THY-peptide (0.5 mg) conjugates were separately emulsified in Freund's complete adjuvant(1:1) and separately injected into SWL rabbits. After three weeks, the rabbits were given an additional booster injections with 0.5 mg of conjugate in incomplete adjuvant. Booster injection were then given with 4-weeks intervals, using the same procedure, up

until totally four booster injections had been given. Sera were collected 12-14 days after the last injection. A specimen of normal rabbits serum (pre-immune serum) was taken from each rabbit before immunization. All sera were aliquoted for storage at -80°C before being used.

## Antibody screening

Cultured cells which were, respectively, expressing and not expressing the MSH receptor, were attached to poly-L-lysine coated slides for 24 hours, and the slides were then gently 10 washed in PBS. (As MSH receptor expressing cells, COS-7 cells transfected with pE-11D, using the method described in Example 3 were used. As controls, which were not expressing MSH receptors, non-transfected COS-7 cells were used). The cells, being attached to the slides, were fixed in 4% 15 paraformaldehyde for 10 minutes at 22°C whereafter the slides were washed twice in PBS. Cells were then permeabilized by incubating in 0.2% Triton X-100 in PBS for 4 minutes at 22°C and the slides were then again washed gently in PBS with 3 changes in 5 minutes intervals. Slides were then pre-incu-20 bated in 10% foetal calf serum for 30 minutes at 22°C whereafter they were incubated with either the pre-immune sera diluted (1:100) or the antisera diluted (1:100) in 10% foetal calf serum, for 60 minutes at 22°C. After this procedure the slides were gently washed in PBS with 3 changes in 5 minute intervals. Slides were then incubated with TRITC-la-25 belled anti-rabbit secondary antibody (diluted 1:40) for 60 minutes at 22°C and then again washed gently in PBS with 3 changes in 5 minute intervals. The cells were then observed under a fluorescent microscope using appropriate filters for the correct wavelengths. 30

## Results:

The cells expressing the MSH receptor showed very little fluorescence when tested with pre-immunesera (reaction can be categorised as +).

The cells expressing the MSH receptor showed high fluorescence both when tested with antisera developed against Peptide M1-Y and when tested with antisera developed against Peptide M2-Y (The reaction can in both cases be categorised as ++++).

The control cells not expressing the MSH receptor showed very little fluorescence when tested with antisera developed against Peptide M1-Y or against Peptide M2-y (the reaction can, in both cases, be categorised as +).

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# SEQUENCE LISTING

(1) GENERAL INFORMATION:	
(i) APPLICANT:	
(A) NAME: Jarl Wikberg	
(B) STREET: Trillvaegen 13	
(C) CITY: Umea	
(E) COUNTRY: Sweden	
(F) POSTAL CODE (ZIP): 905 92 Umea	
(A) NAME: Vijay Chhajlani	
(B) STREET: Stigbergsvæegen	
(C) CITY: Uppsala	
(E) COUNTRY: Sweden	
(F) POSTAL CODE (ZIP): 752 42 Uppsala	
(ii) TITLE OF INVENTION: New polypeptides	
(iii) NUMBER OF SEQUENCES: 16	
(iv) COMPUTER READABLE FORM:	
(A) MEDIUM TYPE: Floppy disk	
(B) COMPUTER: IBM PC compatible	
(C) OPERATING SYSTEM: PC-DOS/MS-DOS	
(D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)	
(2) INFORMATION FOR SEQ ID NO: 1:	
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 1270 base pairs	
(B) TYPE: nucleic acid	
(C) STRANDEDNESS: double	
(D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (cDNA)	
(ix) FEATURE:	
(A) NAME/KEY: CDS	
(B) LOCATION: 1691122	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:	
GGAGAGGGTG TGAGGGCAGA TCTGGGGGTG CCCAGATGGA AGGAGGCAGG CATGGGGGAC	60
ACCCAAGGCC CCCTGGCAGC ACCATGAACT AAGCAGGACA CCTGGAGGGG AAGAACTGTG	120
GGGACCTGGA GGCCTCCAAC GACTCCTTCC TGCTTCCTGG ACAGGACT ATG GCT GTG  Met Ala Val  1	177
net hia vai	

CAG GGA TCC CAG AGA AGA CTT CTG GGC TCC CTC AAC TCC ACC CCC ACA

Gln Gly Ser Gln Arg Arg Leu Leu Gly Ser Leu Asn Ser Thr Pro Thr

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					GGG Gly 25											273
					TCT Ser											321
					GCG Ala											369
AAC Asn	CTG Leu	CAC His 70	TCA Ser	CCC Pro	ATG Met	TAC Tyr	TGC Cys 75	TTC Phe	ATC Ile	TGC Cys	TGC Cys	CTG Leu 80	GCC Ala	TTG Leu	TCG Ser	417
					GGG Gly											465
CTG Leu 100	CTG Leu	GAG Glu	GCC Ala	GGT Gly	GCA Ala 105	CTG Leu	GTG Val	GCC Ala	CGG Arg	GCT Ala 110	GCG Ala	GTG Val	CTG Leu	CAG Gln	CAG Gln 115	513
					GAC Asp											561
					GCC Ala											609
			Arg		CAC His										CGA Arg	657
					TGG Trp		Ala									705
	Ala				CAC His 185	Val					Сув				TTC Phe 195	753
Phe	Leu	Ala	Met	Leu 200		Leu	Met	Ala	Val 205	Leu	Tyr	Val	His	Met 210	Leu	801
Ala	Arg	Ala	Сув 215	Gln	His	Ala	Gln	Gly 220	Ile	Ala	Arg	Leu	His 225	Lys	AGG Arg	849
CAG Gln	CGC	CCG Pro 230	Val	CAC	CAG Gln	GGC Gly	TTT Phe 235	Gly	CTT Leu	AAA Lys	GGC Gly	GCT Ala 240	Val	ACC Thr	CTC Leu	897

				GGC Gly												945
				ATC Ile												993
ATC Ile	TTC Phe	AAG Lys	AAC Asn	TTC Phe 280	AAC Asn	CTC Leu	TTT Phe	CTC Leu	GCC Ala 285	CTC Leu	ATC Ile	ATC Ile	TGC Cys	AAT Asn 290	GCC Ala	1041
ATC Ile	ATC Ile	GAC Asp	CCC Pro 295	CTC Leu	ATC Ile	TAC Tyr	GCC Ala	TTC Phe 300	CAC His	AGC Ser	CAG Gln	GAG Glu	CTC Leu 305	CGC Arg	AGG Arg	1089
			Glu	GTG Val							GCGC	GGT (	GCAC	GCGC'	TT	1139
TAA	GTGT(	GCT (	GGGC.	AGAG	GG A	GGTG:	GTGA'	T AT	TGTG	TGGT	CTG	GTTC	CTG	TGTG:	ACCCT	1199
GGC.	AGTT	CCT	TACC	TCCC	TG G	TCCC	CGTT	T GT	CAAA	GAĢG	ATG	GACT.	AAA	TGAT	CTCTG!	1259
AAG	TGTT	GAA	G													1270
(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:	2:								

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 317 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: polypeptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Met Ala Val Gln Gly Ser Gln Arg Arg Leu Leu Gly Ser Leu Asn Ser

Thr Pro Thr Ala Ile Pro Gln Leu Gly Leu Ala Ala Asn Gln Thr Gly 20 25 30

Ala Arg Cys Leu Glu Val Ser Ile Ser Asp Gly Leu Phe Leu Ser Leu
35 40 45

Gly Leu Val Ser Leu Val Glu Asn Ala Leu Val Val Ala Thr Ile Ala 50 55 60

Lys Asn Arg Asn Leu His Ser Pro Met Tyr Cys Phe Ile Cys Cys Leu
65 70 75 80

Ala Leu Ser Asp Leu Leu Val Ser Gly Ser Asn Val Leu Glu Thr Ala 85 90 95

Val Ile Leu Leu Glu Ala Gly Ala Leu Val Ala Arg Ala Ala Val 100 105 110

- Leu Gln Gln Leu Asp Asn Val Ile Asp Val Ile Thr Cys Ser Ser Met 115 120 125
- Leu Ser Ser Leu Cys Phe Leu Gly Ala Ile Ala Val Asp Arg Tyr Ile 130 135 140
- Ser Ile Phe Tyr Ala Leu Arg Tyr His Ser Ile Val Thr Leu Pro Arg 145 150 155 160
- Ala Arg Arg Val Ala Ala Ile Trp Val Ala Ser Val Val Phe Ser 165 170 175
- Thr Leu Phe Ile Ala Tyr Tyr Asp His Val Ala Val Leu Leu Cys Leu 180 185 190
- Val Val Phe Phe Leu Ala Met Leu Val Leu Met Ala Val Leu Tyr Val 195 200 205
- His Met Leu Ala Arg Ala Cys Gln His Ala Gln Gly Ile Ala Arg Leu 210 215 220
- His Lys Arg Gln Arg Pro Val His Gln Gly Phe Gly Leu Lys Gly Ala 235 240
- Val Thr Leu Thr Ile Leu Leu Gly Ile Phe Phe Leu Cys Trp Gly Pro 245 250 255
- Phe Phe Leu His Leu Thr Leu Ile Val Leu Cys Pro Glu His Pro Thr 260 265 270
- Cys Gly Cys Ile Phe Lys Asn Phe Asn Leu Phe Leu Ala Leu Ile Ile 275 280 285
- Cys Asn Ala Ile Ile Asp Pro Leu Ile Tyr Ala Phe His Ser Gln Glu 290 295 300
- Leu Arg Arg Thr Leu Lys Glu Val Leu Thr Cys Ser Trp 305 310
- (2) INFORMATION FOR SEQ ID NO: 3:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 33 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (synthetic)
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

GGGAATTCTG TGTGTNATCN CNGTGGACCG GTA

(2) INFORMATION FOR SEQ ID NO: 4:

<ul> <li>(i) SEQUENCE CHARACTERISTICS:         <ul> <li>(A) LENGTH: 35 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul> </li> </ul>		
(ii) MOLECULE TYPE: DNA (synthetic)		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:		
GGGGATCCGA AGAAGGGNAA CCAGCAGAGN ATGAA		35
(2) INFORMATION FOR SEQ ID NO: 5:		
(i) SEQUENCE CHARACTERISTICS:		
(A) LENGTH: 285 base pairs (B) TYPE: nucleic acid		
<ul><li>(C) STRANDEDNESS: double</li><li>(D) TOPOLOGY: linear</li></ul>		
(ii) MOLECULE TYPE: DNA (PCR-fragment)		
(ix) FEATURE:		
(A) NAME/KEY: CDS (B) LOCATION: 1285		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:		
TTC TAC GCA CTG CGC TAC CAC AGC ATC GTG ACC ATG CGC		
Phe Tyr Ala Leu Arg Tyr His Ser Ile Val Thr Met Arg 1 5 10	15	
GTG GTG CTT ACG GTC ATC TGG ACG TTC TGC ACG GGG ACT		
Val Val Leu Thr Val Ile Trp Thr Phe Cys Thr Gly Thr 20 25	30	1111
ATG GTG ATC TTC TCC CAT CAT GTG CCC ACA GTG ATC ACC		
Met Val Ile Phe Ser His His Val Pro Thr Val Ile Thr 35 40 45	Phe Ini	per
CTG TTC CCG CTG ATG CTG GTC TTC ATC CTG TGC CTC TAT		
Leu Phe Pro Leu Met Leu Val Phe Ile Leu Cys Leu Tyr 50 55 60	Val H18	met
TTC CTG CTG GCT CGA TCC CAC ACC AGG AAG ATC TCC ACC	CTC CCC	AGA 240
Phe Leu Leu Ala Arg Ser His Thr Arg Lys Ile Ser Thr 65 70 75	Leu Pro	80
GCC AAC ATG AAA GGG GCC ATC ACC CTC ACC ATC CTG CTG	GGC ATT	285
Ala Asn Met Lys Gly Ala Ile Thr Leu Thr Ile Leu Leu 85 90	95	

J 74/ U	40/4															
									1	00						
(2)	INFO	ORMAT	NOI	FOR	SEQ	ID N	10: 6	<b>:</b>								
	(	(	A) LE 3) TY	NGTH	CHAR H: 95 amin GY:	ami	ino a									
	(ii)	MOI	LECUI	E TY	PE:	poly	pept	ide								
	( <b>x</b> i)	SE	QUENC	E DI	ESCRI	PTIC	ON: 5	SEQ I	D NO	): 6:	•					
Phe 1	Tyr	Ala	Leu	Arg 5	Tyr	His	Ser	Ile	Val 10	Thr	Met	Arg	Arg	Thr 15	Val	
Val	Val	Leu	Thr 20	Val	Ile	Trp	Thr	Phe 25	Cys	Thr	Gly	Thr	Gly 30	Ile	Thr	
Met	Val	Ile 35	Phe	Ser	His	His	Val 40	Pro	Thr	Val	Ile	Thr 45	Phe	Thr	Ser	
Leu	Phe 50	Pro	Leu	Met	Leu	Val 55	Phe	Ile	Leu	Сув	Leu 60	Tyr	Val	His	Met	
Phe 65	Leu	Leu	Ala	Arg	Ser 70	His	Thr	Arg	Lys	Ile 75	Ser	Thr	Leu	Pro	Arg 80	
Ala	Asn	Met	Lys	Gly 85	Ala	Ile	Thr	Leu	Thr 90	Ile	Leu	Leu	Gly	Ile 95		
(2)		) SE (	QUEN A) L B) T	CE C ENGT YPE:	SEQ HARA H: 3 nuc DEDN	CTER 06 b leic	ISTI ase aci	CS: pair d	g							
		•	•		OGY:											
	(ii	) MO	LECU	LE T	YPE:	DNA	(PC	R-fr	agme	nt)						
	(ix	(	-	AME/	KEY:											
	(xi	.) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0: 7	:					
TTC Phe	TAC	GCA Ala	CTG	CGC	TAC	CAC	AGC Ser	ATC	GTG Val	ACG Thr	GCG Ala	AGG Arg	CGC	TCA Ser	GGG	•

				CGC Arg 5												48
GCC Ala	ATC Ile	ATC Ile	GCC Ala 20	GGC Gly	ATC Ile	TGG Trp	GCT Ala	TTC Phe 25	TGC Cys	ACG Thr	GGC Gly	TGC Cys	GGC Gly 30	ATT Ile	GTC Val	96
TTC Phe	ATC Ile	CTG Leu 35	TAC Tyr	TCA Ser	GAA Glu	TCC Ser	ACC Thr 40	TAC Tyr	GTC Val	ATC Ile	CTG Leu	TGC Cys 45	CTC Leu	ATC Ile	TCC Ser	144

ATG Met	TTC Phe 50	TTC Phe	GCT Ala	ATG Met	CTG Leu	TTC Phe 55	CTC Leu	CTG Leu	GTG Val	TCT Ser	CTG Leu 60	TAC Tyr	ATA Ile	CAC His	ATG Met	192
TTC Phe 65	CTC Leu	CTG Leu	GCG Ala	CGG Arg	ACT Thr 70	CAC His	GTC Val	AAG Lys	CGG Arg	ATC Ile 75	GCG Ala	CTC	TGC Cys	CCG Pro	GGG 80	240
CCA Pro	GCT Ala	CTG Leu	CGC <b>A</b> rg	GGC Gly 85	AGA Arg	GGA Gly	CCA Pro	GCA Ala	TGC Cys 90	AGG Arg	GGC Gly	GCG Ala	GTC Val	ACC Thr 95	CTC Leu	288
			CTG Leu 100													306

- (2) INFORMATION FOR SEQ ID NO: 8:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 102 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: polypeptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

Phe Tyr Ala Leu Arg Tyr His Ser Ile Val Thr Ala Arg Arg Ser Gly
1 5 10 15

Ala Ile Ile Ala Gly Ile Trp Ala Phe Cys Thr Gly Cys Gly Ile Val 20 25 30

Phe Ile Leu Tyr Ser Glu Ser Thr Tyr Val Ile Leu Cys Leu Ile Ser 35 40 45

Met Phe Phe Ala Met Leu Phe Leu Leu Val Ser Leu Tyr Ile His Met 50 55 60

Phe Leu Leu Ala Arg Thr His Val Lys Arg Ile Ala Leu Cys Pro Gly 65 70 75 80

Pro Ala Leu Arg Gly Arg Gly Pro Ala Cys Arg Gly Ala Val Thr Leu 85 90 95

Thr Ile Leu Leu Gly Ile 100

- (2) INFORMATION FOR SEQ ID NO: 9:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 312 base pairs
    - . (B) TYPE: nucleic acid
      - (C) STRANDEDNESS: double
      - (D) TOPOLOGY: linear

	(ii)	MOI	ECUL	E TY	PE:	DNA	(PCR	-fra	gmen	t)						
	(ix)	(2	TURE () NA () LC	ME/K			12									
	(xi)	SEÇ	QUENC	E DE	SCRI	PTIC	N: S	EQ I	D NC	: 9:						
TTC Phe	TAC Tyr	GCA Ala	CTG Leu	CGT Arg 5	TAC Tyr	CAC His	AGC Ser	ATC Ile	GTG Val 10	ACC Thr	GTG Val	CGG Arg	CGG Arg	GCC Ala 15	CTC Leu	48
ACC Thr	TTG Leu	ATC Ile	GTG Val 20	GCC Ala	ATC Ile	TGG Trp	GTC Val	TGC Cys 25	TGC Cys	GGC Gly	GTC Val	TGT Cys	GGC Gly 30	GTG Val	GTG Val	96
TTC Phe	ATC Ile	GTC Val 35	TAC Tyr	TCG Ser	GAG Glu	AGC Ser	AAA Lys 40	ATG Met	GTC Val	ATT Ile	GTG Val	TGC Cys 45	CTC Leu	ATC Ile	ACC Thr	144
ATG Met	TTC Phe 50	Phe	GCC Ala	ATG Met	ATG Met	CTC Leu 55	CTC Leu	ATG Met	GGC Gly	ACC	CTC Leu 60	TAC Tyr	GTG Val	CAC His	ATG Met	192
TTC Phe 65	Leu	TTT Phe	GCG Ala	CGG	CTG Leu 70	CAC	GTC Val	AAG Lys	CGC Arg	ATA Ile 75	GCA Ala	GCA Ala	CTG Leu	CCA Pro	CCT Pro 80	240
GCC	GAC	GGG Gly	GTG Val	GCC Ala 85	Pro	CAG Gln	CAA Gln	CAC His	TCA Ser 90	Сув	ATG Met	AAG Lys	GGG	GCA Ala 95	GTC Val	288
			ATC Ile 100	Leu							,					312
(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:	10:			•					
` ,		(i)	SEQU (A) L (B) I	ENCE ENGT	CHA	RACI 04 a	ERIS mino cid	TICS								
	(13	L) MO	DLECU	ILE 1	YPE:	pol	yper	tide	2							
	(x:	L) SI	EQUE	ICE I	ESCF	RIPTI	ON:	SEQ	ID N	10: 1	LO:					
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			20	)				2!	5				3(	J	l Val	
Ph	e Il	e Va		r Se	r Glu	ı Se	r Lyı		t Va	l Ile	e Vai	1 Cy	s Le	ı Ile	• Thr	

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Met	Phe 50	Phe	Ala	Met	Met	Leu 55	Leu	Met	Gly	Thr	Leu 60	Tyr	Val	His	Met	
Phe 65	Leu	Phe	Ala	Arg	Leu 70	His	Val	Lys	Arg	Ile 75	Ala	Ala	Leu	Pro	Pro 80	
Ala	Asp	Gly	Val	Ala 85	Pro	Gln	Gln	His	Ser 90	Сув	Met	Lys	Gly	<b>Ala</b> 95	Val	
Thr	Leu	Thr	Ile 100	Leu	Leu	Gly	Ile									
(2)	INFO	RMAT	ION	FOR	SEQ	ID N	io: 1	11:								
	(i)	SEC	UENC	E CH	IARAC	TERI	STIC	cs:								
	, ,	(P	) LE	engti	1: 37	12 ba	rae I	pairs	3							
		•	•		nucl											
		•	•		EDNE CGY:			ote								
		(1	, 10	POLC	ŊΙ.	TTILE	<b>.a.</b>									
	(ii)	MOI	ECUI	E T	PE:	DNA	(PCI	R-fra	gmer	it)						
		. 50	mirot	D 4												
	(IX)	FE?			KEY:	CDS										
		-	•	-	ON:		372									
	( <b>x</b> i)	) SEÇ	UENC	CE DI	escr:	[PTI	ON:	SEQ :	ID NO	): 11	l:					
OMC.	<b>m</b> c m	GTG	እጥ <b>ር</b>	GCG	CTG	GAC	CGG	TAC	ATC	TCC	ATC	TTC	TAC	GCA	CTG	48
		Val														
1	•			5					10					15		
CGC	TAC	CAC	AGC	ATC	GTG	ACC	CTG	CCG	CGG	GCG	CCG	GAA	GCC	GTT	GCG	96
		His														
_	_		20					25					30			
000	እጥሮ	TGG	CTC	GCC	AGT	GTC	GTC	TTC	AGC	ACG	CTC	TTC	ATC	GCC	TAC	144
Ala	Ile	Trp	Val	Ala	Ser	Val	Val	Phe	Ser	Thr	Leu	Phe	Ile	Ala	Tyr	
		35					40					45				
<b>ma</b> 0					<b>6</b> 55	OMC.	OTC.	TOO	СТС	CTG	GTC	TTC	TTC	CTG	GCT	192
TAC	~~~															
Tur	GAC	CAC	GTG Val	Ala	Val	Leu	Leu	Cys	Leu	Val	Val	Phe	Phe	Leu	Ala	
Tyr	GAC Asp 50	His	GTG Val	Ala	Val	Leu 55	Leu	Сув	Leu	Val	Val	Phe	Phe	Leu	Ala	
Tyr	<b>Asp</b> 50	His	Val	Ala	Val	Leu 55	Leu	Сув	Leu	Val	Val	Phe	Phe	Leu	Ala	240
Tyr ATG	Asp 50 CTG	His GTG	Val	Ala	Val GCC	Leu 55	Leu	Cys TAC	Leu : GTC	Val	Val 60 ATG	Phe	Phe	Leu	Ala	240
Tyr ATG	Asp 50 CTG	His	Val	Ala	Val GCC	Leu 55 GTG Val	Leu	Cys TAC	Leu : GTC	Val	Val 60 ATG Met	Phe	Phe	Leu	Ala	240
Tyr ATG Met	Asp 50 CTG Leu	His GTG Val	Val CTC Leu	Ala ATG Met	GCC Ala	Leu 55 GTG Val	CTG Leu	TAC Tyr	Leu GTC Val	CAC His	Val 60 ATG Met	Phe CTG Leu	GCC Ala	CGG Arg	GCC Ala 80	
Tyr ATG Met	Asp 50 CTG Leu	His GTG Val	Val CTC Leu	Ala ATG Met	Val GCC Ala 70	Leu 55 GTG Val	CTG Leu	TAC Tyr	Leu GTC Val	CAC His 75	Val 60 ATG Met	Phe CTG Leu	GCC Ala	CGC Arg	GCC Ala 80 CCG	240 288
Tyr ATG Met	Asp 50 CTG Leu	His GTG Val	Val CTC Leu	Ala ATG Met	GCC Ala 70 GGC	Leu 55 GTG Val	CTG Leu	TAC Tyr	Leu GTC Val	CAC His 75 CAC	Val 60 ATG Met	Phe CTG Leu	GCC Ala	CGC Arg	GCC Ala 80 CCG Pro	

GTC CAC CAG GGC TTT GGC CTT AAA GGC GCT GTC ACC CTC ACC ATC CTG

Val His Gln Gly Phe Gly Leu Lys Gly Ala Val Thr Leu Thr Ile Leu

100

CTG GGC ATT TTC ACC GTC TCG TGG CGC CCC TTC TTC Leu Gly Ile Phe Thr Val Ser Trp Arg Pro Phe Phe 115

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- (2) INFORMATION FOR SEQ ID NO: 12:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 124 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

Leu Cys Val Ile Ala Leu Asp Arg Tyr Ile Ser Ile Phe Tyr Ala Leu 1 5 10 15

Arg Tyr His Ser Ile Val Thr Leu Pro Arg Ala Pro Glu Ala Val Ala 20 25 30

Ala Ile Trp Val Ala Ser Val Val Phe Ser Thr Leu Phe Ile Ala Tyr 35 40 45

Tyr Asp His Val Ala Val Leu Leu Cys Leu Val Val Phe Phe Leu Ala 50 55 60

Met Leu Val Leu Met Ala Val Leu Tyr Val His Met Leu Ala Arg Ala 65 70 75 80

Cys Gln His Ala Gln Gly Ile Ala Arg Leu His Lys Arg Gln Arg Pro

Val His Gln Gly Phe Gly Leu Lys Gly Ala Val Thr Leu Thr Ile Leu 100 105 110

Leu Gly Ile Phe Thr Val Ser Trp Arg Pro Phe Phe 115

- (2) INFORMATION FOR SEQ ID NO: 13:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 35 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: double
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

GGGAATTCTA CGCACTGCGC TACCACAGCA TCGTG

(2) INFORMATION FOR SEQ ID NO: 14:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:         <ul> <li>(A) LENGTH: 34 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul> </li> </ul>	
(ii) MOLECULE TYPE: DNA (synthetic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
GGGGATCCAA TGCCCAGCAG GATGGTGAGG GTGA	34
(2) INFORMATION FOR SEQ ID NO: 15:	
<ul> <li>(i) SEQUENCE CHARACTERISTICS:         <ul> <li>(A) LENGTH: 1650 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: double</li> <li>(D) TOPOLOGY: linear</li> </ul> </li> </ul>	
(ii) MOLECULE TYPE: DNA (cDNA)	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 6161590	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	
CTTGAGGAGA ATGTCGTGCA GTAGCCTTAG GAATGTGAAC ATTGGGAGAC TGGCTGGGAT	60
TTTGTAGGTT ATGAGAAGGG GACACTTATG ATATGTGAAC TTGAGCCCAG GAGAGAAGCC	120
ATAAAAAGTG AAACTGTCCT GGGCACTTGG AGGTGAGTGT CTCTCTAGTA AGATGCATGT	180
GAAAGGCCTG GGAGCTGAAA GCAAGGAGAG CAGAAGAGGC TGGTGAAGAT TCTAATCTGC	240
GTGTCCAGGG GCACTCTTCC AGGTCTCAGG AACGCAGGTC AGAATGTGCA AGCCAGCTGC	300
CGGGCACGTG GCTCACCCCT GTAGTACCAG CACTTTGGGA GGCTGAGAGA GAAGATCGCT	360
TGTGGCCAGG AGTTTGAGAC CAGACTGGGG CTTCATAGGG AGACCCTGTC TCTTAAAAAA	420
AAAAAAAAA AAGGACTGAG TGAGCCGAGC CCAGTCCTCT CATGCACTGT GTCATTCATC	480
CCCTTTCTTA GGCTGTGTTG GTTCTAGGCT AGCTGCTGTC TTTCTTTGGT AGGCTGCTAA	540
CCTCTTTGGA TTGTGAATTT AAAACATGTT TTACAGTAAA TTTGCTGCCA AGACAAGAGG	600
TGTATTTCTC CAGCA ATG AAT TCC TCA TTT CAC CTG CAT TTC TTG GAT CTC  Met Asn Ser Ser Phe His Leu His Phe Leu Asp Leu  1 5 10	651
AAC CTG AAT GCC ACA GAG GGC AAC CTT TCA GGA CCC AAT GTC AAA AAC	699

Asn Leu Asn Ala Thr Glu Gly Asn Leu Ser Gly Pro Asn Val Lys Asn

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AAG Lys	TCT Ser 30	TCA Ser	CCA Pro	TGT Cys	GAA Glu	GAC Asp 35	ATG Met	GGC Gly	ATT Ile	GCT Ala	GTG Val 40	GAG Glu	GTG Val	TTT Phe	CTC Leu	747
ACT Thr 45	CTG Leu	GGT Gly	GTC Val	ATC Ile	AGC Ser 50	CTC Leu	TTG Leu	GAG Glu	AAC Asn	ATC Ile 55	TTG Leu	GTC Val	ATA Ile	GGG Gly	GCC Ala 60	795
ATA Ile	GTG Val	AAG Lys	AAC Asn	AAA Lys 65	AAC Asn	CTG Leu	CAC His	TCC Ser	CCC Pro 70	ATG Met	TAC Tyr	TTC Phe	TTC Phe	GTG Val 75	TGC Cys	843
AGC Ser	CTG Leu	GCA Ala	GTG Val 80	GCG Ala	Asp GAC	ATG Met	CTG Leu	GTG Val 85	AGC Ser	ATG Met	TCC	AGT Ser	GCC Ala 90	TGG Trp	GAG Glu	891
ACC Thr	ATC Ile	ACC Thr 95	ATC Ile	TAC Tyr	CTA Leu	CTC Leu	AAC Asn 100	AAC ABn	AAG Lys	CAC His	CTA Leu	GTG Val 105	ATA Ile	GCA Ala	GAC Asp	939
GCC Ala	TTT Phe 110	Val	CGC Arg	CAC His	ATT Ile	GAC Asp 115	AAT Asn	GTG Val	TTT Phe	GAC Asp	TCC Ser 120	ATG Met	ATC Ile	TGC Cys	ATT Ile	987
TCC Ser 125	Val	GTG Val	GCA Ala	TCC Ser	ATG Met 130	TGC Cys	AGC Ser	TTA Leu	CTG Leu	GCC Ala 135	ATT	GCA Ala	GTG Val	GAT Asp	AGG Arg 140	1035
TAC Tyr	GTC Val	ACC Thr	ATC	TTC Phe 145	Tyr	GCC Ala	CTG Leu	CGC Arg	TAC Tyr 150	His	CAC	ATC	ATG Met	ACG Thr 155	GCG Ala	1083
<b>A</b> GG <b>A</b> rg	CGC Arg	TCA Ser	GGG Gly 160	Ala	ATC	ATC	GCC Ala	GGC Gly 165	Ile	TGG Trp	GCT	TTC Phe	TGC Cys 170	Thr	GGC Gly	1131
TGC Cys	GGC Gly	ATT	Val	TTC	ATC	CTG Leu	TAC Tyr 180	Ser	GAA Glu	TCC Ser	ACC Thr	TAC Tyr 185	Val	ATC	CTG Leu	1179
TGC Cys	CTC Leu 190	Ile	TCC Ser	ATG Met	TTC Phe	TTC Phe 195	Ala	ATG Met	CTG Leu	TTC Phe	Leu 200	Leu	GTG Val	TCT Ser	CTG Leu	1227
TAC Ty: 20:	: Ile	CAC His	C ATG	TTC Phe	CTC Leu 210	Leu	G GCG	CGG Arg	ACT Thr	CAC His 215	Val	C AAG L Lys	CGG Arg	ATC Ile	GCG Ala 220	1275
CT( Lev	C TGC	C CCC	G GGG	9 CC2 7 Pro 225	Ala	CTC	Arg	G GG(	AGA Arg 230	Gly	CCI Pro	A GCA	TGG Tr	GAG Glr 235	GGC Gly	1323
GC6 Ala	G GTG	C ACC	C GT( r Val 240	L Thi	C ATO	CTO	G CTO	G GGG 1 Gly 24	y Val	G TT	T AC	c GTC	TG0 L Cys 250	Tr	GCC Ala	1371

			CTT Leu			Thr										1419
			CGC Arg							Met						1467
			TCC Ser						Ile							1515
GAG Glu	ATG Met	CGG Arg	AAG Lys	ACC Thr 305	TTT Phe	AAG Lys	GAG Glu	ATT	ATT Ile 310	TGC Cys	TGC Cys	CGT Arg	GGT Gly	TTC Phe 315	AGG Arg	1563
			AGC Ser 320						TAAC	GACA	AA (	TGCI	CCTC	T		1610
CTGT	rggc:	rct (	STTCI	CCTI	T GI	TTGC	TCAC	CTA	TGAC	AAA						1650
(2)	INF	ORMA!	rion	FOR	SEQ	ID N	10: 1	16:								
			SEQUE A) LI													
		(I	B) T													
		(1	ጋነ ጥር	DOI.	CY:	line										
	, , , ,	•	D) TO				ar									
		) <b>MO</b> :	LECUI	LE T	YPE:	prot	ear cein	SFO '	TD NO	n• 1 <i>6</i>	<b>.</b>					
	(xi	) MO	OUEN(	LE T	YPE: ESCR	prot	ear cein					<b>h</b> an	Leu	Aan	λla	
Met 1	(xi	) MO	LECUI	LE T	YPE: ESCR	prot	ear cein					Asn	Leu	Asn 15	Ala	
1	(xi Asn	) MO	OUEN(	CE DI Phe	YPE: ESCR: His	prot	ear cein ON:	Phe	Leu 10	Asp	Leu			15		
1 Thr	(xi Asn Glu	) MO: ) SE Ser	DUENC Ser Asn 20 Met	CE Di Phe 5	YPE: ESCR: His	prot IPTIC Leu Gly	ear cein ON:	Phe Asn 25	Leu 10 Val	Asp Lys	Leu	Lys	Ser 30	15 Ser	Pro	
Thr Cys	(xi Asn Glu Ser 50	) MO: Ser Gly Asp 35	LECUI QUENC Ser Asn 20 Met	Phe 5 Leu Gly	YPE: ESCR: His Ser Ile	prof IPTIC Leu Gly Ala Ile 55	ear ein ON: His Pro Val 40	Phe Asn 25 Glu Val	Leu 10 Val Val	Asp Lys Phe Gly	Leu Asn Ala 60	Thr 45	Ser 30 Leu Val	Ser Gly Lys	Pro Val Asn	
Thr Cys Ile	(xi Asn Glu Ser 50 Asn	) MO: ) SE Ser Gly Asp 35	QUENC Ser Asn 20 Met	CE Di Phe 5 Leu Gly Glu	YPE: ESCR: His Ser Ile Asn Pro	prof IPTIC Leu Gly Ala Ile 55	ear cein ON: His Pro Val 40 Leu Tyr	Phe Asn 25 Glu Val	Leu 10 Val Val Ile	Asp Lys Phe Gly Val	Leu Asn Leu Ala 60 Cys	Thr 45 Ile	Ser 30 Leu Val	Ser Gly Lys	Pro Val Asn Val 80	
Thr Cys Ile Lys 65	(xi Asn Glu Ser 50 Asn	) MO: ) SE Ser Gly Asp 35 Leu Met	DUENG Ser Asn 20 Met Leu His	CE Di Phe 5 Leu Gly Glu Ser	YPE: ESCR: His Ser Ile Asn Pro 70 Ser	prof IPTIC Leu Gly Ala Ile 55 Met	ein ON: His Pro Val 40 Leu Tyr	Asn 25 Glu Val Phe	Leu 10 Val Val Ile Phe Ala 90	Asp Lys Phe Gly Val 75	Leu Asn Leu Ala 60 Cys	Thr 45 Ile Ser	Ser 30 Leu Val	Ser Gly Lys Ala Thr	Pro Val Asn Val 80	
Thr Cys Lys 65 Ala	Clu Ser 50 Asr	Ser  Gly  Asp  35  Leu  Met	DUENG Ser Asn 20 Met Leu His	CE Di Phe 5 Leu Gly Ser Val 85	YPE: ESCR: His Ser Ile Asn Pro 70 Ser	Prof IPTIC Leu Gly Ala Ile 55 Met	ear ein ON: His Pro Val 40 Leu Tyr	Phe Asn 25 Glu Val Phe Ser Val 105	Leu 10 Val Val Ile Phe Ala 90	Asp Lys Phe Gly Val 75 Trp	Leu Ala 60 Cys Glu Asp	Thr 45 Ile Ser	Ser 30 Leu Val Leu Ile	Ser Gly Lys Ala Thr 95 Val	Pro Val Asn Val 80	

Ser Met Cys Ser Leu Leu Ala Ile Ala Val Asp Arg Tyr Val Thr Ile 130 135 140

Phe Tyr Ala Leu Arg Tyr His His Ile Met Thr Ala Arg Arg Ser Gly
145 150 155 160

Ala Ile Ile Ala Gly Ile Trp Ala Phe Cys Thr Gly Cys Gly Ile Val 165 170 175

Phe Ile Leu Tyr Ser Glu Ser Thr Tyr Val Ile Leu Cys Leu Ile Ser 180 185 190

Met Phe Phe Ala Met Leu Phe Leu Leu Val Ser Leu Tyr Ile His Met 195 200 205

Phe Leu Leu Ala Arg Thr His Val Lys Arg Ile Ala Leu Cys Pro Gly 210 215. 220

Pro Ala Leu Arg Gly Arg Gly Pro Ala Trp Gln Gly Ala Val Thr Val 225 230 235 240

Thr Met Leu Leu Gly Val Phe Thr Val Cys Trp Ala Pro Phe Phe Leu 245 250 255

His Leu Thr Leu Met Leu Ser Cys Pro Gln Asn Leu Tyr Cys Ser Arg 260 265 270

Phe Met Ser His Phe Asn Met Tyr Leu Ile Leu Ile Met Cys Asn Ser 275 280 285

Val Met Asp Pro Leu Ile Tyr Ala Phe Arg Ser Gln Glu Met Arg Lys 290 295 300

Thr Phe Lys Glu Ile Ile Cys Cys Arg Gly Phe Arg Ile Ala Cys Ser 305 310 315

Phe Pro Arg Arg Asp 325

- (2) INFORMATION FOR SEQ ID NO: 17:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 22 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (synthetic)
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO 17:

GGAAGCTTTC TTTGGTAGGC TG

(2)	INFORMATION FOR SEQ ID NO: 18:	
	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 22 base pairs</li><li>(B) TYPE: nucleic acid</li></ul>	
	(C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:	
GGT	CTAGAGC CACAGAGAGG AG	22
(2)	INFORMATION FOR SEQ ID NO: 19:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 17 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:	
CTG	CATTTCT TGGATCT	17
(2)	INFORMATION FOR SEQ ID NO: 20:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 17 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (synthetic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:	
AAG	CTGCACA TGGATGC	17

## INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism reference on page 37, line 10	rred to in the description
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet X
Name of depositary institution  Deutsche Sammlung von Mikroorgani	ismen und Zellkulturen GmbH (DSM)
Address of depositary institution (including postal code and country)  Mascheroder Weg 1b  D-38124 Braunschweig  Federal Republic of Germany	
Date of deposit 9 August 1993	Accession Number DSM 8440
C. ADDITIONAL INDICATIONS (leave blank if not applicable	(c) This information is continued on an additional sheet
As regards the respective Patent designated states, the applicant deposited microorganisms only be nominated by the requester until is granted or the date on which fused or withdrawn or is deemed	made available to an expert the date on which the patent the application has been re-
D. DESIGNATED STATES FOR WHICH INDICATION	ONS ARE MADE (if the indications are not for all designated States)
E. SEPARATE FURNISHING OF INDICATIONS (leav	re blank if not applicable)
	Burcau later (specify the general nature of the indications e.g., "Accession
Engagining Office use only	For International Bureau use only
For receiving Office use only  This sheet was received with the international application	
Authorized officer Susamixfedunce	Authorized officer

Form PCT/RO/134 (July 1992)

## INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism reference on page 36, line 27	erred to in the description
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution  Deutsche Sammlung von Mikroorgan:	ismen und Zellkulturen GmbH (DSM)
Address of depositary institution (including postal code and country)  Mascheroder Weg lb D-38124 Braunschweig Federal Republic of Germany	
Date of deposit 24 August 1992	Accession Number DSM 7214
C. ADDITIONAL INDICATIONS (leave blank if not applicab	ole) This information is continued on an additional sheet
As regards the respective Patent designated states, the applicant deposited microorganisms only be nominated by the requester until is granted or the date on which fused or withdrawn or is deemed	made available to an expert the date on which the patent the application has been re-
D. DESIGNATED STATES FOR WINCH INDICATE	
E. SEPARATE FURNISHING OF INDICATIONS (lea	ve blank if not applicable)
The indications listed below will be submitted to the International Number of Deposit*)	al Bureau later (specify the general nature of the indications e.g., "Accession
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	This sheet was received by the International Bureau on:
Authorized officer  Susanny feldmun	Authorized officer

Form PCT/RO/134 (July 1992)

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#### CLAIMS

- 1. A DNA fragment having the nucleotide sequence shown in SEQ ID NO: 1 or an analogue or subsequence thereof which
  - 1) has a homology with the DNA sequence shown in SEQ ID NO: 1 of at least 50%, and/or
    - 2) encodes a polypeptide, the amino acid sequence of which is at least 50% homologous with the amino acid sequence shown in SEQ ID NO: 2, and/or
- 3) encodes a polypeptide which binds an antibody which is also bound by an MSH receptor, and/or
  - 4) encodes a polypeptide which is an MSH receptor or which has the same binding capacity as an MSH receptor.
- 2. A DNA fragment having the nucleotide sequence shown in SEQ ID NO: 15 or an analogue or subsequence thereof which
  - 1) has a homology with the DNA sequence shown in SEQ ID NO: 15 of at least 50%, and/or
- 2) encodes a polypeptide, the amino acid sequence of which is at least 50% homologous with the amino acid sequence shown in SEQ ID NO: 16, and/or
  - 3) encodes a polypeptide which binds an antibody which is also bound by an MSH receptor, and/or
  - 4) encodes a polypeptide which is an MSH receptor or which has the same binding capacity as an MSH receptor.
  - 3. A DNA fragment which is a subsequence of the DNA fragment according to claim 1 or 2 which comprises at least 15 nucleo-

tides, preferably at least 18 nucleotides, more preferably at least 21 nucleotides, even more preferably at least 27 nucleotides and most preferably at least 51 nucleotides.

- 4. A DNA fragment according to claim 1 encoding a polypeptide comprising amino acids no. 1-317 shown in SEQ ID NO: 2.
  - 5. A DNA fragment according to claim 2 encoding a polypeptide comprising amino acids no. 1-325 shown in SEQ ID NO: 16.
- 6. A DNA fragment which shows at least 55% homology, preferably at least 70%, more preferably at least 80% and most preferably at least 95% homology, to the DNA sequence shown in SEQ ID NO: 1 or SEQ ID NO: 15.
  - 7. A DNA fragment which is a modified DNA fragment or a subsequence thereof which differs from the DNA sequence SEQ ID NO: 1, SEQ ID NO: 15, SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9 or the corresponding subsequence in that at least one nucleotide has been substituted, added, inserted, deleted and/or rearranged.
- 8. A DNA fragment which is a fusion DNA fragment comprising a DNA fragment according to any of the claims 1-7 in frame with one or more second DNA fragments different from or identical to the DNA fragment according to any of claims 1-7, the second DNA fragment preferably selected from the group consisting of DNA fragments encoding, diphtheria toxin, a staphylococcus protein, a ricin toxin, a pseudomonas endotoxin, abrin and fungal ribosome-inactivation proteins (RIP), the resulting DNA fragment encoding a fusion protein.

#### 9. A DNA fragment having

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the nucleotide sequence shown in SEQ ID NO: 3 or an analogue thereof, wherein the nucleotides 13 and/or 15 and/or 23 optionally are substituted by C, or

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the nucleotide sequence shown in SEQ ID NO: 4, or an analogue thereof, wherein the nucleotides 19 and/or 29 and/or 32 optionally are substituted by C and wherein the nucleotides 20 and/or 31 optionally are substituted by G.

- 10. A polypeptide having the amino acid sequence shown in SEQ ID NO: 2 or an analogue or subsequence thereof which
  - 1) is an MSH receptor or is capable of binding to MSH or an analogue thereof, and/or
- 2) is encoded by a DNA fragment which is at least 50% homologous with the DNA fragment shown in SEQ ID NO: 1, and/or
  - 3) binds an antibody which is also bound by an MSH receptor.
- 15 11. A polypeptide having the amino acid sequence shown in SEQ ID NO: 16 or an analogue or subsequence thereof which
  - 1) is an MSH receptor or is capable of binding to MSH or an analogue thereof, and/or
- 2) is encoded by a DNA fragment which is at least 50% homologous with the DNA fragment shown in SEQ ID NO: 15, and/or
  - 3) binds an antibody which is also bound by an MSH receptor.
- 12. A polypeptide which is a subsequence of the polypeptide
  25 according to claim 10 comprising from 5 to 316 amino acids,
  preferably at least 7 amino acids, more preferably at least
  10 amino acids, even more preferably at least 15 amino acids
  and most preferably at least 30 amino acids.

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- 13. A polypeptide which is a subsequence of the polypeptide according to claim 11 comprising from 5 to 324 amino acids, preferably at least 7 amino acids, more preferably at least 10 amino acids, even more preferably at least 15 amino acids and most preferably at least 30 amino acids.
- 14. A polypeptide which shows at least 55% homology, preferably at least 70%, more preferably at least 80% and most preferably at least 95% homology, to the polypeptide shown in SEQ ID NO: 2 or the polypeptide shown in SEQ ID NO: 16.
- 10 15. A polypeptide consisting of or comprising

a subsequence of the polypeptide shown in SEQ ID NO: 2, the subsequence being selected from the group consisting of amino acids 1-40, 99-117, 181-189, 268-277, 62-76, 141-158, 212-244, 300-317, 39-63, 75-100, 116-141, 157-182, 188-213, 243-269 and 276-301, and analogues thereof, or

a subsequence of the polypeptide shown in SEQ ID NO: 16 the subsequence being selected from the group consisting of amino acids 1-38, 97-115, 179-187, 265-274, 61-74, 138-156, 211-240, 297-326, 37-62, 73-98, 114-139, 155-180, 186-212, 239-266 and 273-298, and analogues thereof.

- 16. A polypeptide according to any of claims 10-15 which is glycosylated and/or coupled to a carbohydrate or lipid moiety and/or contains a palmitoyl anchor or a part thereof and/or provided with a detectable label and/or coupled to a solid support.
- 17. A polypeptide which is a fusion polypeptide comprising a polypeptide according to any of claims 10-16 or a subsequence thereof fused to a second polypeptide which may be different from or identical to the polypeptide according to any of

claims 10-16, which fusion polypeptide preferably has retained the capability of binding to MSH or an analogue thereof.

- 18. A polypeptide according to any of claims 10-17 in substantially pure form.
- 5 19. A polypeptide according to any of claims 10-18 in lipid soluble form.
  - 20. A DNA fragment or a subsequence or analogue thereof which
- shows a homology with any of the nucleotide sequences
  shown in SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9 of
  at least 40%, preferably 50%, more preferably at least
  55%, even more preferably at least 70%, still more
  preferably at least 80% and most preferably at least
  95%, and/or
- which can be isolated by using the nucleotide sequence shown in SEQ ID NO: 13 and/or SEQ ID NO: 14 as a primer, and/or

which has the nucleotide sequence shown in SEQ ID NO: 5, SEQ ID NO: 7 or SEQ ID NO: 9.

- 20 21. A DNA fragment which is a fusion DNA fragment comprising a DNA fragment according to claim 20 in frame with one or more second DNA fragments different from or identical to the DNA fragment according to claim 20, the second DNA fragment preferably being selected from the group consisting of DNA fragments encoding a melanotropic hormone receptor, an MSH receptor and an ACTH receptor.
- 22. A polypeptide or a subsequence or analogue thereof, which shows a homology of at least 40%, preferably at least 50%, more preferably at least 55%, even more preferably at least 70%, still more preferably at least 80% and most preferably

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at least 95% with the polypeptide shown in SEQ ID NO: 6, SEQ ID NO: 8 or SEQ ID NO: 10.

- 23. A polypeptide which is a fusion polypeptide comprising a polypeptide according to claim 22 or a subsequence thereof fused to a second polypeptide which may be different from or identical to the polypeptide according to claim 22, the second polypeptide preferably being selected from the group consisting of a melanotropic hormone receptor, an MSH receptor and an ACTH receptor or an analogue or subsequence thereof.
  - 24. A DNA fragment coding for a polypeptide as defined in any of claims 10-19, 22 and 23.
- 25. A replicable expression vector carrying a DNA fragment according to any of claims 1-9, 20, 21 and 24, which vector is capable of replicating in a host organism or a cell line.
  - 26. A vector according to claim 25 which is

pB-11D deposited under the deposition number DSM 7214 at Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH, or

- pE-MC-2 deposited under the deposition number DSM 8440 at Deutsche Sammlung von Mikroorganismen und Zellkultu-ren GmbH.
- 27. A cell which carries and is capable of replicating the DNA fragment according to any of the claims 1-9, 20, 21 and 25.
  - 28. A cell according to claim 27, which is selected from the group consisting of a bacterium, a yeast and a protozoan, or the cell is derived from a multicellular organism selected from the group consisting of a fungus, an insect, a plant, and a mammal, and the cell preferably being a bacterium

selected from the group consisting of the genus Bacillus, Escherichia and Salmonella.

- 29. A method of producing a polypeptide as defined in any of claims 10-19, 22 and 23, comprising the following steps of:
- 5 (a) inserting a DNA fragment as defined in any of the claims 1-9, 20, 21 and 24 into an expression vector,
- (b) transforming a suitable host cell according to claim 27 or 28 with the vector produced in step(a),
  - (c) cultivating the host cell produced in step (b) under suitable conditions for expressing the polypeptide,
  - (d) harvesting the polypeptide, and
- (e) optionally subjecting the polypeptide to posttranslational modification,

or comprising liquid and/or solid phase peptide synthesis procedures.

- 30. A stable cell line which produces the polypeptide according to claim 10 or 11 and which preferably contains and expresses cDNA encoding the polypeptide of claim 10 or claim 11.
- 31. A method of preventing or stimulating the coupling of an MSH receptor to its guanine nucleotide binding protein in an animal, in particular a mammal, comprising administering a substance which in advance has been found to bind to a polypeptide according to any of claims 10-19 so as to occupy one or several of the cytoplasmic loops and/or the C-terminal sequence.

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32. A method of preventing or stimulating the binding of MSH and similar peptides to an MSH receptor in an animal, in particular a human, comprising administering, to the animal, a substance which in advance has been found to bind to a polypeptide according to any of claims 10-19 so as to occupy the binding site of the receptor using an antagonist, a blocker or a compound such as a derivative of MSH having a structure similar to MSH, and optionally thereby preventing or stimulating the generation of second messenger elements.

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- 33. A method of increasing or decreasing the generation of second messenger elements, and/or increasing or decreasing the production of an MSH receptor and/or optionally increasing or decreasing the binding affinity of MSH to an MSH receptor, comprising administering to an animal, in particular a human, a medicament which is or becomes bound to a substance which in advance has been found to bind to a polypeptide according to any of claims 10-19.
- 34. A method of targeting, with a medicament, a cell that contains an MSH receptor on its surface, comprising admini20 stering a substance optionally linked to a medicament which substance in advance has been found to bind to a polypeptide according to any of claims 10-19 and which substance binds to the MSH receptor.
- 35. A method according to the claim 34 wherein the medicament is a radionuclide or a toxin or any other molecule of natural or synthetic origin.
- 36. A method according to any of claims 31-35 for the treatment of an MSH receptor expressing disease condition selected from the group consisting of melanoma, skin cancer, vitiligo, pyretic condition, inflammatory condition, nociceptive condition, catatonic condition, impaired memory condition, reduced or increased skin tanning, pigmentation condition, epilepsy and nerve damage.

- 37. A method for treating conditions caused by MSH receptor deficiency or impaired MSH receptor function in a mammal, such as a human, comprising administering a polypeptide according to claim 19 which is an MSH receptor, or is capable of binding to MSH, or is an analogue thereof, to the mammal.
- 38. A method for treating conditions caused by MSH receptor deficiency or impaired MSH receptor function, such as tyrosinase-positive albinism, in a mammal, such as a human, comprising introducing a DNA fragment according to any of claims 1-9, 20, 21 and 24 encoding an active form of an MSH receptor.
- 39. A method for increasing or decreasing the melanin content of the skin in a mammal, such as a human, comprising administering substances as defined in any of claims 31-34 that are active through an MSH receptor, preferably to increase the skin tanning without or with reduced exposure to sunlight or to avoid sunburns.
- 40. A method of activating the antipyretic and/or anti-inflammatory and/or antinociceptive and/or memory improving
  and/or nerve regenerating action effected via an MSH receptor, comprising administering a substance that acts on the
  MSH receptor to bring about the antipyretic and/or antiinflammatory and/or antinociceptive and/or memory improving
  and/or nerve regenerating actions.
- 25 41. A method of diagnosing an MSH receptor expressing disease condition such as melanoma or skin cancer, comprising targeting a cell containing an MSH receptor on its surface with a diagnostic agent capable of binding to the MSH receptor, which diagnostic agent can be detected following binding to the receptor.
  - 42. The method according to claim 41, as used in the assessment of the prognosis and/or guidance for further treatment of melanoma or skin cancer.

- 43. A method for detecting an MSH receptor in a biological sample, such as a tissue sample, a cell culture or a cell suspension, wherein the sample is treated with a optionally labelled substance that binds to the MSH receptor, and detecting or visualizing the presence of the bound substance.
  - 44. An antibody optionally provided with a detectable label which antibody is reactive with a polypeptide according to any of claims 10-19, 22 and 23 and which is preferably a monoclonal antibody.
- 10 45. A method for detection and/or quantitation of the MSH receptor mRNA, comprising extracting RNA from a biological sample such as a cell, a tissue sample, a cell culture or a cell suspension and measuring the hybridization of said RNA to a labelled DNA fragment according to any of claims 1-9, 20, 21 and 24 or a labelled RNA fragment constructed from the DNA fragment according to any of claims 1-9, 20, 21 and 24.
  - 46. A method for detection and/or quantitation of the MSH receptor mRNA, comprising extracting RNA from cells or tissues and converting it into cDNA for subsequent use in the polymerase chain reaction (PCR), preferably using PCR primer(s) which is/are synthesized based on the DNA fragment claimed in any of claims 1-9, 20, 21 and 24.
- 47. The use of any of the methods of claims 43, 45 and 46 for diagnosis of an MSH receptor expressing disease condition such as melanoma and skin cancer.
  - 48. The use of a DNA fragment according to any of claims 1-9, 20, 21 and 24 for the isolation of other similar DNA fragments using techniques such as PCR or hybridization.
- 49. The use of a polypeptide according to any of claims 10-19, 22 and 23 for designing DNA probes for use in techniques such as PCR and hybridization.

- 50. The use of a polypeptide according to any of claims 1019, 22 and 23 for the deduction of the three-dimensional
  structure of an MSH receptor or an analogue thereof having
  MSH binding capacity for use in the design of a substance
  capable of binding to the MSH receptor.
- 51. A method for selecting a substance which is capable of binding to a melanotropic hormone receptor polypeptide such as an MSH receptor and which substance may optionally be capable of preventing or stimulating the generation of a second messenger element in a cell such as a mammalian cell, in particular a human cell, by its binding to the melanotropic hormone receptor polypeptide the method comprising one or more of the following steps:
- 1a) incubating a sample containing a melanotropic hormone
  receptor polypeptide or an analogue thereof, the melanotropic hormone receptor polypeptide preferably being a
  polypeptide according to any of claims 10-19, 22 and 23,
  with radioactively labelled melanotropic hormone or an
  analogue thereof and with the substance to be tested,
  and
- 1b) measuring the binding affinity of the substance to be tested for the melanotropic hormone receptor polypeptide by separating bound from free labelled melanotropic hormone or an analogue thereof using either filtration, centrifugation, superflow or chromatography followed by measuring the radioactivity retained in the sample by standard nuclear counting,

or

incubating a sample containing a melanotropic hormone receptor polypeptide or an analogue thereof, the melanotropic hormone receptor polypeptide preferably being a polypeptide according to any of

claims 10-19, 22 and 23, with melanotropic hormone or an analogue thereof and with the substance to be tested, and

- measuring the binding affinity of the substance to 2b) be tested for the melanotropic hormone receptor 5 polypeptide by separating free melanotropic hormone or the analogue thereof from the bound melanotropic hormone or the analogue thereof using either filtration, centrifugation, superflow or chromatography followed by measuring the bound melanotropic 10 hormone or the analogue thereof by a detection system capable of detecting melanotropic hormone or the analogue thereof, preferably using a detection system such as radio immunoassay, immunefluorescense assay, UV light absorption spectrometry or fluo-15 rescence emission spectrometry.
- 52. A method for selecting a substance which is—capable of binding to a melanotropic hormone receptor polypeptide such as an MSH receptor and which substance may optionally be capable of preventing or stimulating the generation of a second messenger element in a cell such as a mammalian cell, in particular a human cell, by its binding to the melanotropic hormone receptor polypeptide the method comprising one or more of the following steps:
- incubating a sample containing a melanotropic hormone receptor polypeptide or an analogue thereof linked to a solid support, the melanotropic hormone receptor polypeptide preferably being a polypeptide according to any of claims 10-19, 22 and 23, with melanotropic hormone or an analogue thereof and with the substance to be tested, and
  - 1b) measuring the binding affinity of the substance to be tested for the melanotropic hormone receptor polypeptide or an analogue thereof by separating

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free melanotropic hormone or an analogue thereof from the melanotropic hormone or an analogue thereof bound to the melanotropic hormone receptor polypeptide or analogue thereof by washing, followed by measuring the bound melanotropic hormone or an analogue thereof by using a ligand, preferably an antibody, capable of binding to the bound melanotropic hormone or an analogue thereof which ligand is in itself detectable, or which ligand is a first ligand which can be rendered detectable using a second ligand, preferably an antibody capable of binding to the said first ligand,

or

- incubating a sample containing, preferably in a soluble form or in a solid phase being attached to a matrix, the melanotropic hormone receptor polypeptide which preferably is a polypeptide according to any of claims 10-19, 22 and 23 or an analogue thereof, with melanotropic hormone and with the substance to be tested, and
  - 2b) measuring the alteration in the degree of interaction of the melanotropic hormone receptor with a G-protein caused by the binding of the substance to be tested to the melanotropic hormone receptor.
- 25 53. A method according to any of claims 31-34, 39, 40, 43, and 50-52 wherein the substance is an antibody or a part thereof or a molecule of natural or synthetic origin having affinity for an MSH receptor or the melanotropic hormone receptor polypeptide.

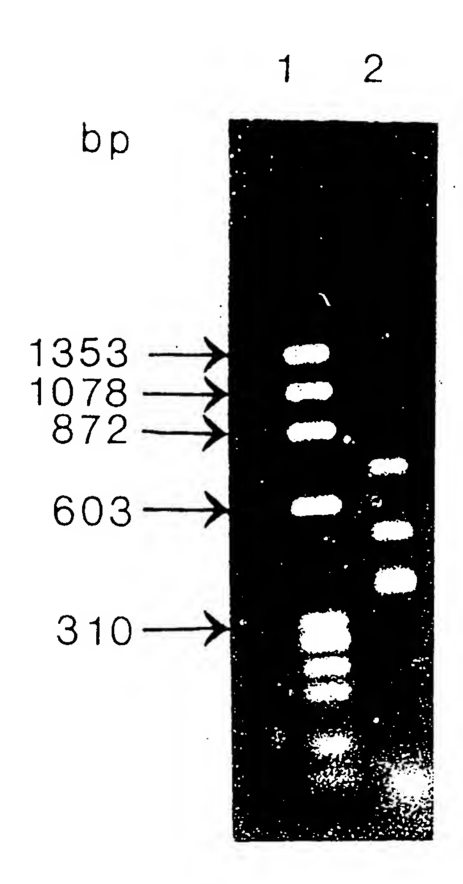


Fig. 1

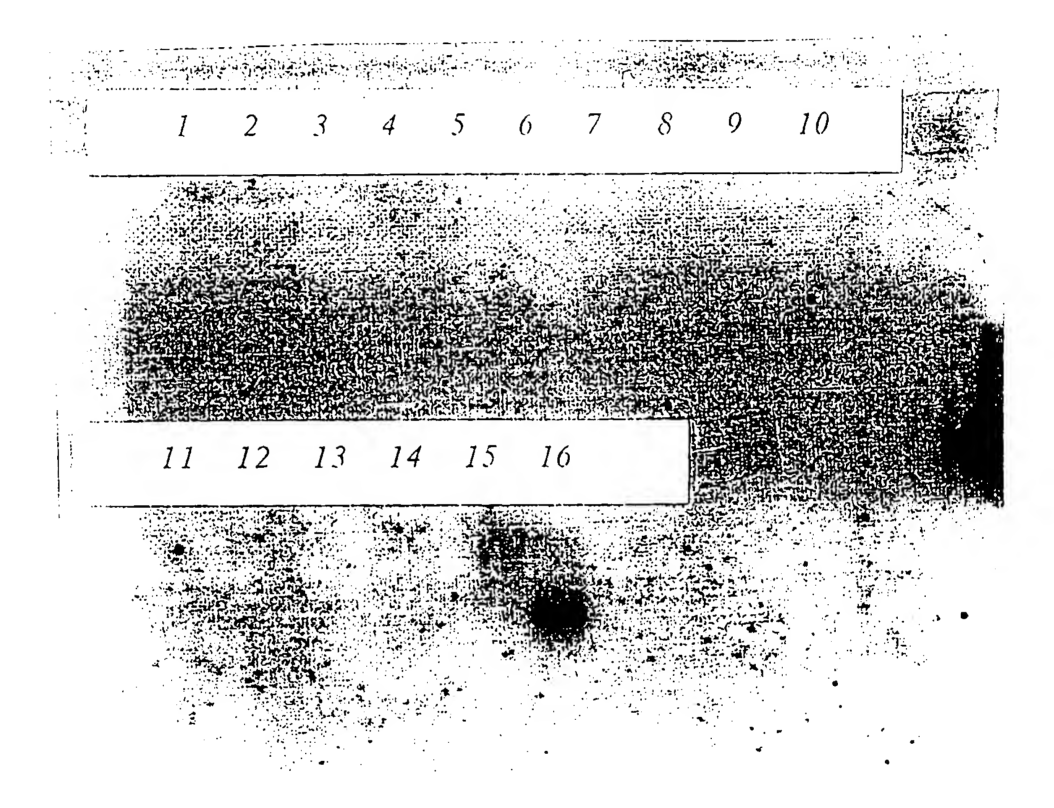


Fig. 2

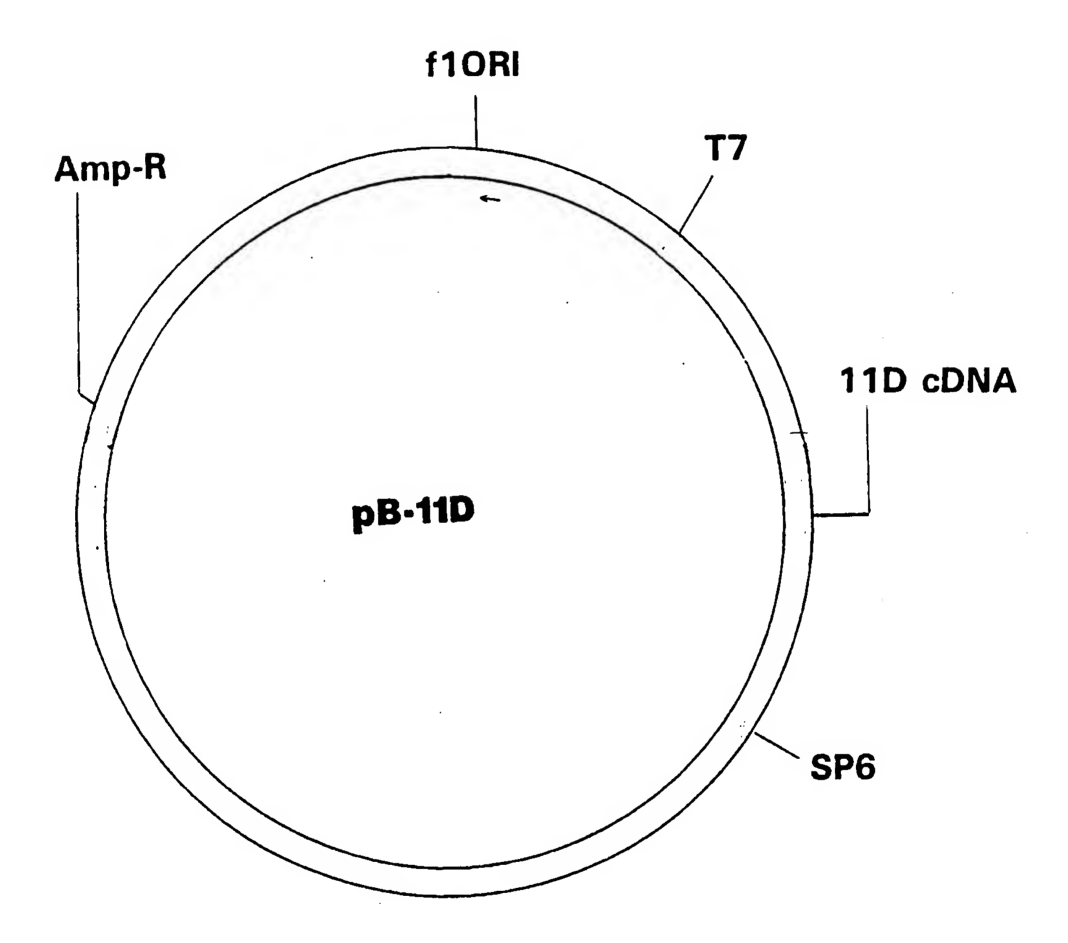


Fig. 3

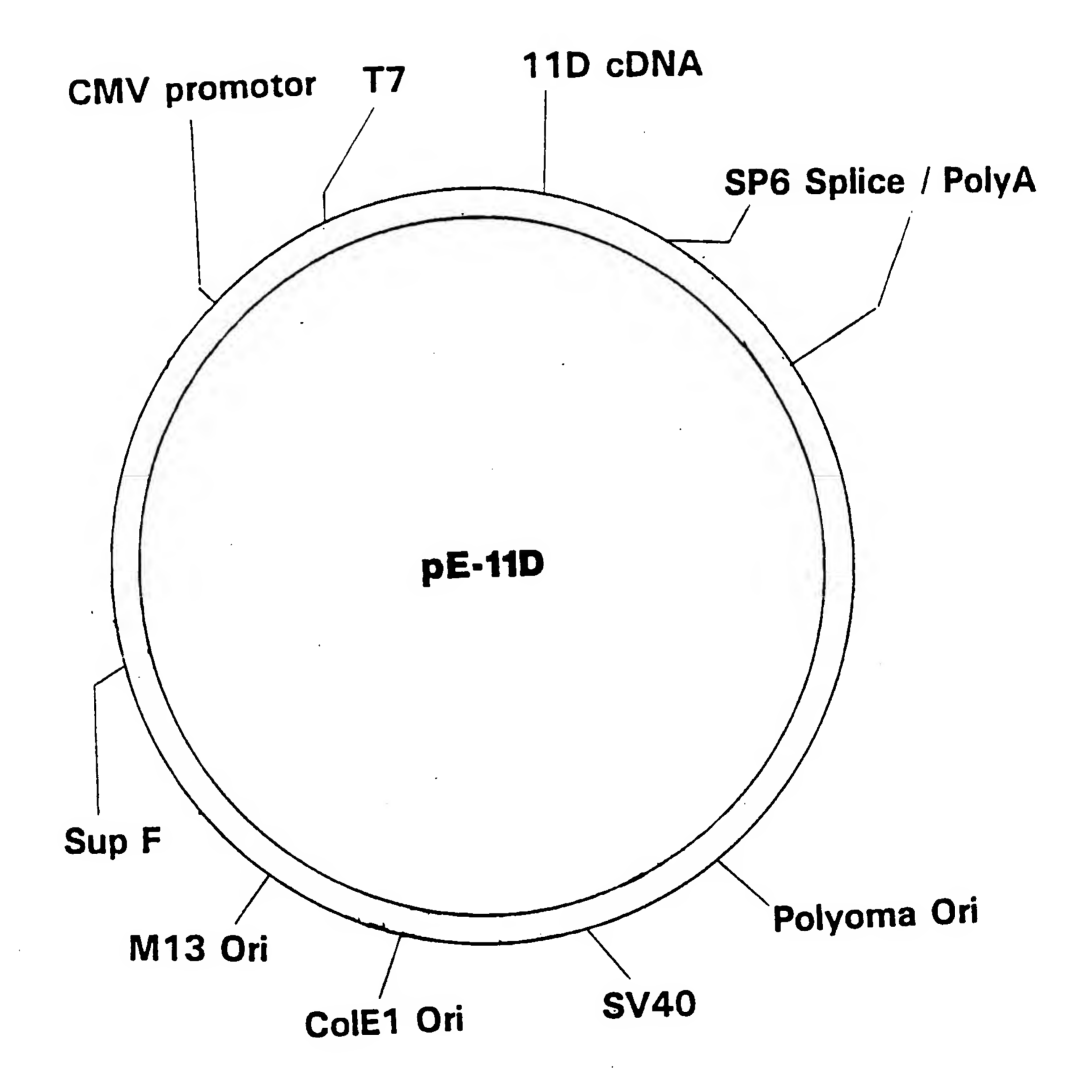


Fig. 4

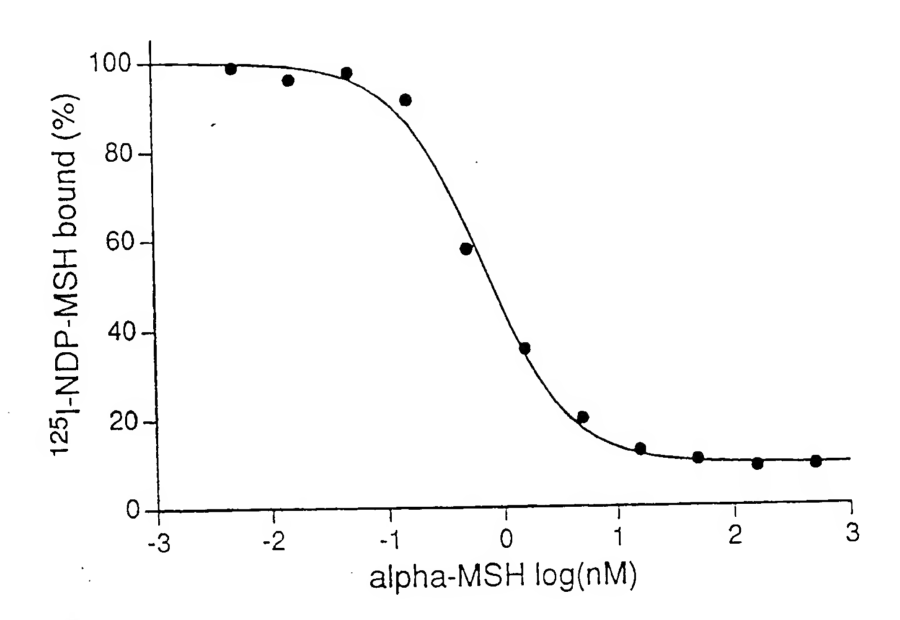


Fig. 5

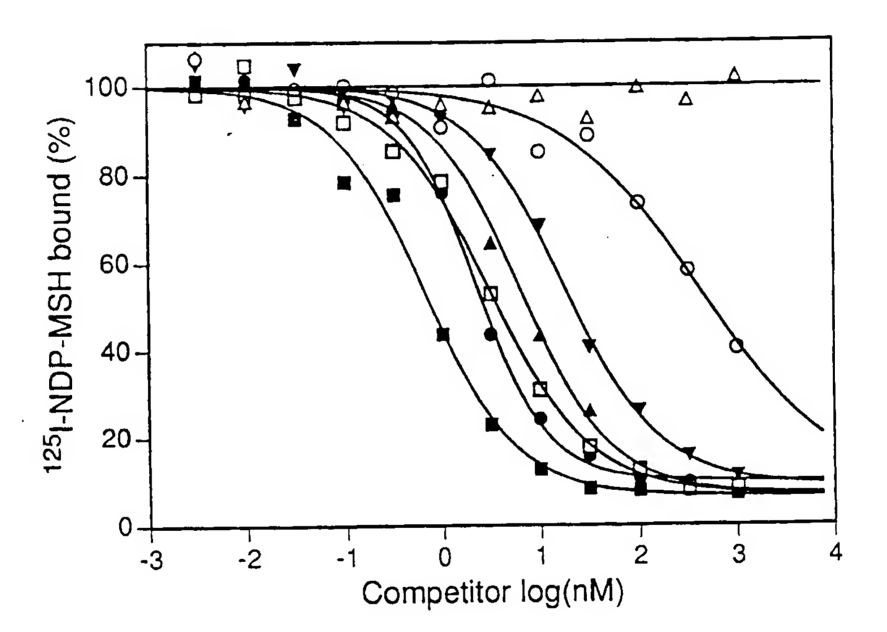


Fig. 6

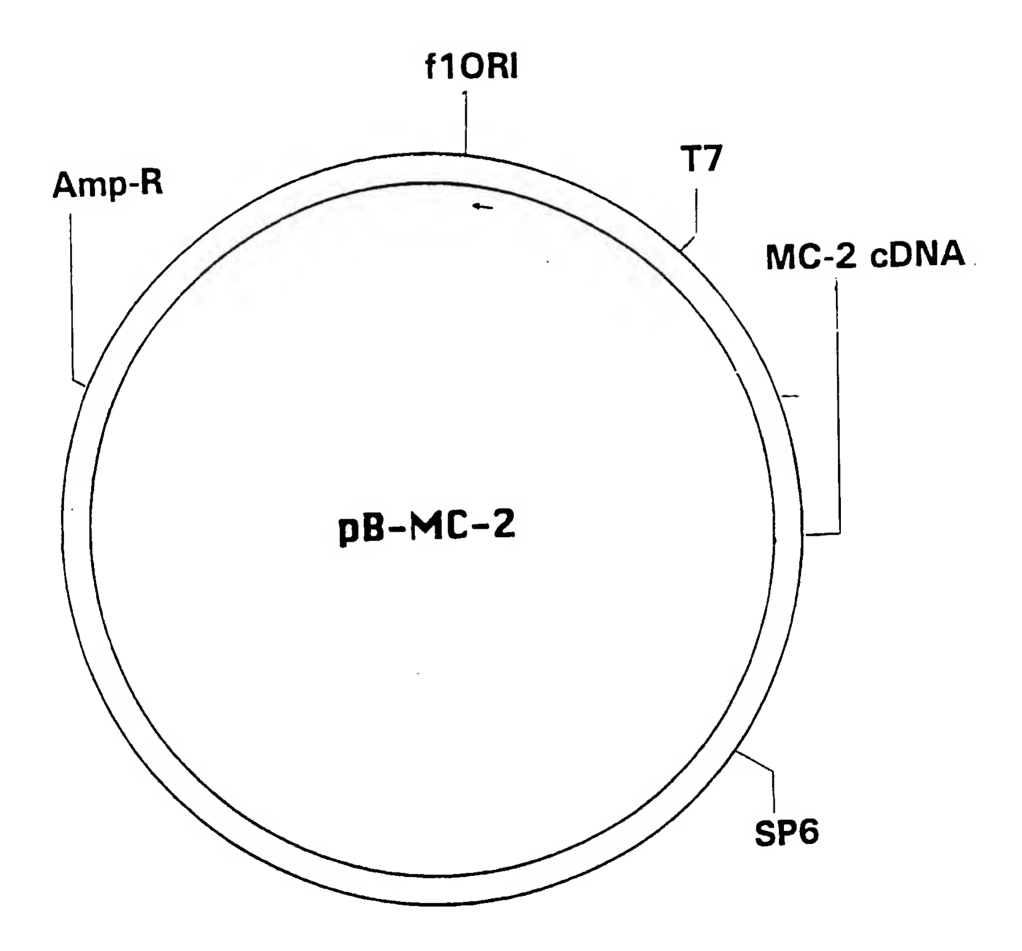


Fig. 7

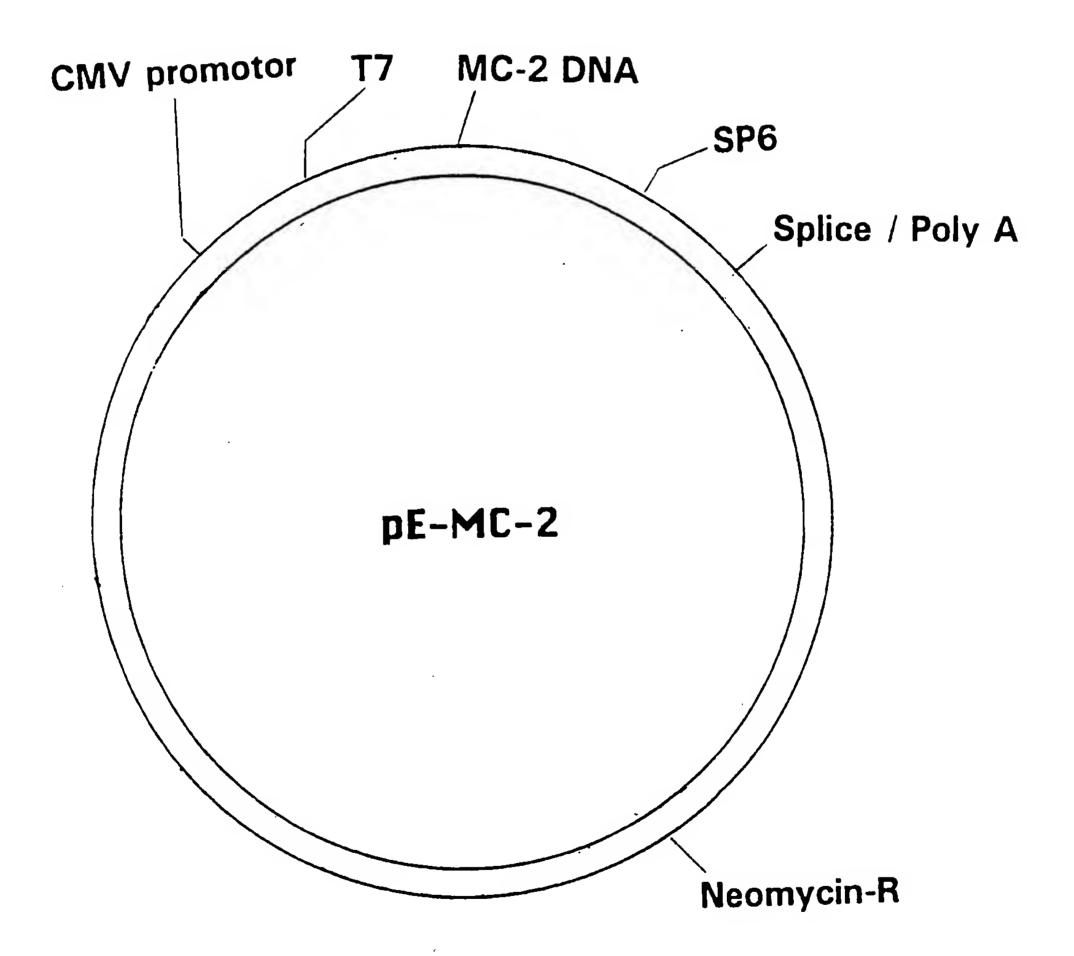


Fig. 8

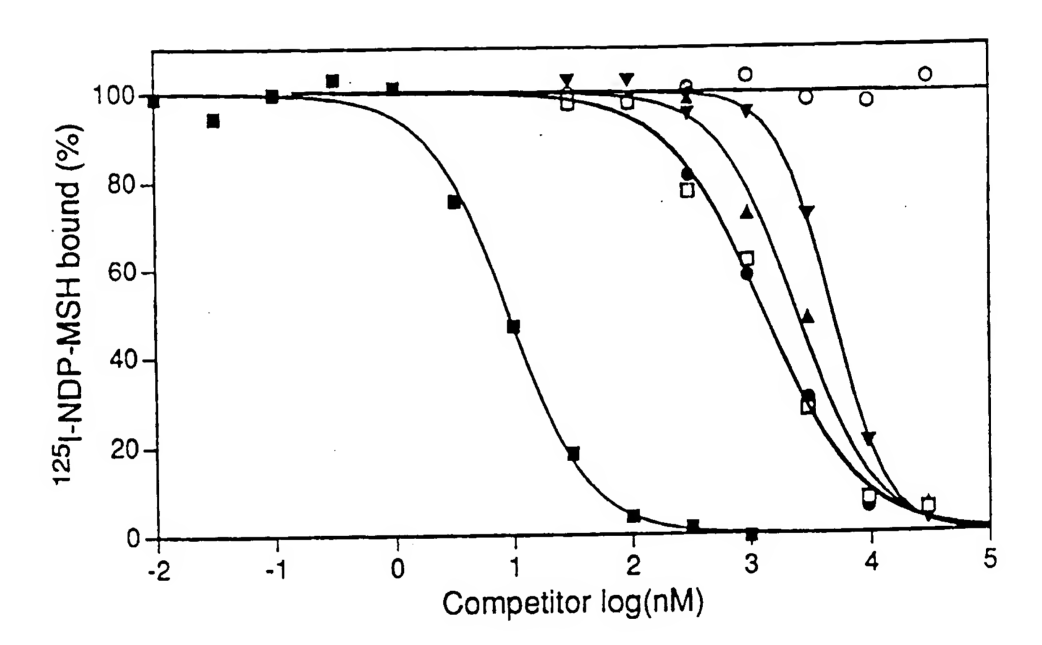


Fig. 9

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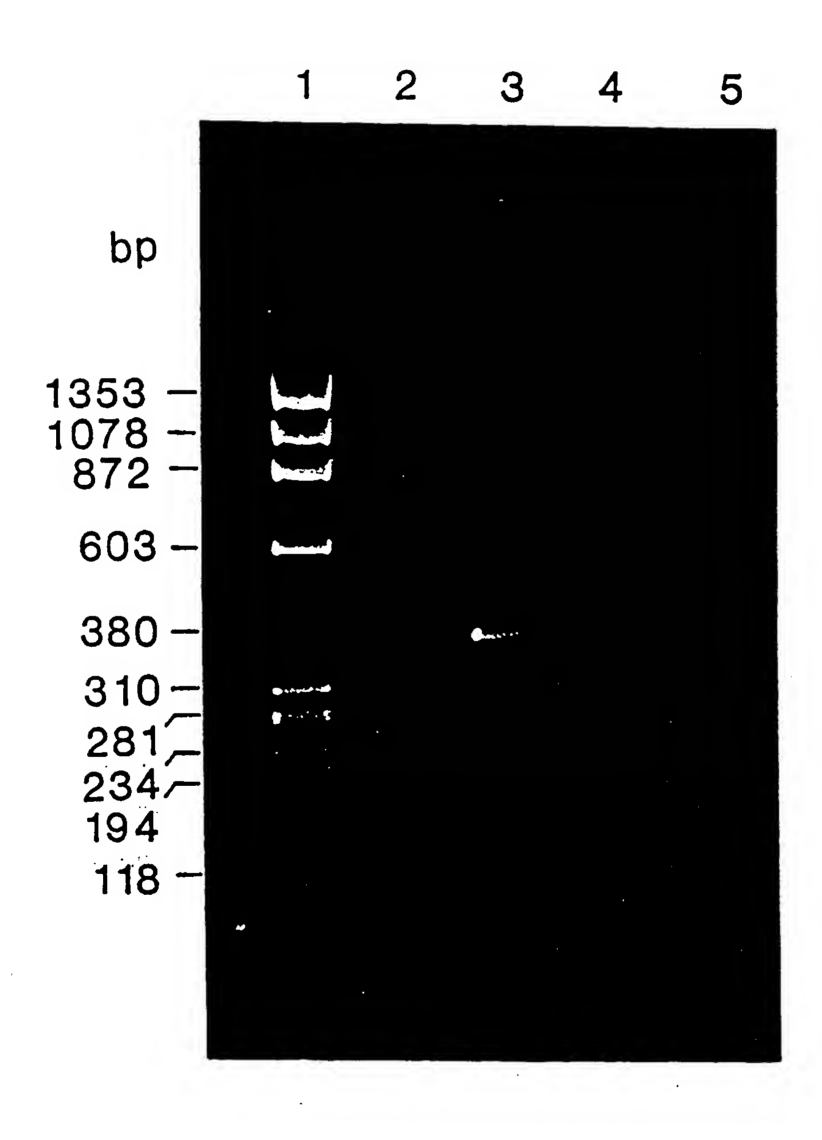


Fig. 10

### INTERNATIONAL SEARCH REPORT

Internetional Application No PCT/DK 93/00273

A. CLASS IPC 5	C12N15/12 C12N15/62 C07K13 C12P21/08 G01N33/52	/00 A61K37/02	G01N33/53	
According	to International Patent Classification (IPC) or to both national cl	assification and IPC	<del></del> <del></del>	
	S SEARCHED	inches or mhole)		
IPC 5	documentation searched (classification system followed by classification s	ication symbols)		
Document	ation searched other than minimum documentation to the extent the	hat such documents are included in t	the fields searched	
Electronic	data base consulted during the international search (name of data	hase and, where practical, search to	erms used)	
C. DOCUI	MENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of the	ne relevant passages	Relevant to claim No.	
Ρ,Χ	SCIENCE vol. 257 , 28 August 1992 , LAN US pages 1248 - 1251		1,3,4,6, 10,18, 25-30	
	KATHLEEN G. MOUNTJOY ET AL. 'The of a family of genes that encode melanocortin receptors.' see the whole document	de the		
P,X	FEBS LETTERS vol. 309, no. 3, 14 September AMSTERDAM NL pages 417 - 420 YIJAY CHHAJLANI ; JARL E.WIKBERG cloning and expression of the melanocyte stimulating hormone cDNA.	G 'Molecular human	1,3,4,6, 10,18, 25-30	
	see the whole document			
		-/		
X Fu	orther documents are listed in the continuation of box C.	Patent family member	s are listed in annex.	
* Special of	categories of cited documents:  ment defining the general state of the art which is not	"T" later document published a	after the international filing date a conflict with the application but inciple or theory underlying the	
considered to be of particular relevance  "E" earlier document but published on or after the international filing date  "L" document which may throw doubts on priority claim(s) or  "L" document which may throw doubts on priority claim(s) or				
whice citate of the citate of	th is cited to establish the publication date of another tion or other special reason (as specified)  Iment referring to an oral disclosure, use, exhibition or imeans	cannot be considered to it document is combined wi	levance; the claimed invention nvolve an inventive step when the th one or more other such docu- being obvious to a person skilled	
'P' docu	ment published prior to the international filing date but r than the priority date claimed	'&' document member of the	والمساور والمساور والموازم والمساور والمساور والمساور والمساور والمساور والمساور والمساور	
Date of the	he actual completion of the international search	Date of mailing of the inte		
	7 December 1993	06-0	- 1994	
Name an	d mailing address of the ISA  European Patent Office, P.B. 5818 Patentlaan 2	Authorized officer		
<u> </u>	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+31-70) 340-3016	Delanghe, l	_	

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### INTERNATIONAL SEARCH REPORT

International Application No PCT/DK 93/00273

∐Continu		PC1/UK 93/UU2/3
	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
P,X	JOURNAL OF BIOLOGICAL CHEMISTRY vol. 268, no. 20 , 15 July 1993 , BALTIMORE, MD US pages 15174 - 15179 IRA GANTZ ET AL. 'Molecular cloning ,expression, and gene localization of a fourth melanocortin receptor.' see the whole document	1,3,4,6, 10,18, 25-30
P, X	JOURNAL OF BIOLOGICAL CHEMISTRY vol. 268, no. 11 , 15 April 1993 , BALTIMORE, MD US pages 8246 - 8250 IRA GANTZ ET AL. 'Molecular cloning of a novel melanocortin receptor.' see the whole document	1,3,4,6, 10,18, 25-30

International application No.

### INTERNATIONAL SEARCH REPORT

PCT/DK93/00273

Bux 1	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
Flus inte	rnational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
2.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:  Remark: Although claims 31-40 completely and 41-42 partially as far as the they eman on "in vivo" method, are directed to a method of treatment or diagnostic method practised on the human/animal body the search has been carried out and based on the alleged effects of the compound/composition.  Claims Nos.:  because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
	Claims Nos.; because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inter	national Scarching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all earchable claims.
2	As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3	As only some of the required additional search fees were timely paid by the applicant, this international search report overs only those claims for which fees were paid, specifically claims Nos.:
4. N	to required additional search fees were timely paid by the applicant. Consequently, this international search report is estricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark "	The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.